COMMERCIAL ORGANIC ANALYSIS;

A TREATISE ON THE

PROPERTIES, PROXIMATE ANALYTICAL EXAMINATION, AND MODES OF ASSAYING THE VARIOUS ORGANIC CHEMICALS AND PRODUCTS EMPLOYED IN THE ARTS, MANUFACTURES, MEDICINE, &c.;

WITH CONCISE METHODS FOR

THE DETECTION AND DETERMINATION OF THEIR IMPURITIES, ADULTERATIONS, AND PRODUCTS OF DECOMPOSITION

37

ALFRED H ALLEN, FI.C. FCS.

PAST PERSIDENT OF THE SOCIETY OF PUBLIC ANALYSTS, PUBLIC ANALYST FOR THE WEST MIDNO OF TORKSHEER, THE NORTHERN DIVISION OF DEEDISHEE THE COUNTY-BOROUGH OF SERFFEELD, &C

Second Edition, Rebised and Enlarged

VOLUME III-PART II

(Reprinted 1899)

AMINES AND AMMONIUM BASES, HYDRAZINES, BASES FRUM TAR, VEGETABLE ALKALUIDS.

LONDON

J & A CHURCHILL

7 GREAT MARLEOROUGH STREET



It is ten years since the publication of the last edition of that part of Commercial Organic Analysis which treated of Alkaloids and Tar Basss. These subjects then occupied about 120 pages. In the edition now issued 570 pages have already been printed, and I feel reductantly compelled to publish the subject-matter now ready as Part II. of Volume III., leaving the sections on the less important Alkaloids and the chapters on Animal Basss, Cyangeen Compounds, PROTEIDS, &c., to be issued separated as Part III.

In Part II., now published, I have endeavoured to describe fully and accurately such of the Organic Bases as have any practical interest, and to give reliable information as to their sources. The AMINES. HYDRAZINES, and PYRIDINE and its Derivatives are now considered for the first time. The ANTIPYRETICS, and other synthetical remedies with which modern Chemistry has enriched medicine, are described fully, in cases where they fall appropriately within the scope of the present Volume, and I believe the sections on Antipyrine, Antifebrin, Phenacetin, Thalline, &c., contain a resumé of all published information on their respective subjects In the Chapter on Vegetable ALKALOIDS I have spared no pains to render the more important articles as complete and trustworthy as possible, and in this endeavour have received most valuable assistance from Mr W Chattaway, Mr A J Cownley, Mr R A Crupps, Mr D B Dott, Mr A W Gerrard, Mr O. Helmer, Dr B. H Paul, Mr M J Sheridan, Dr C R. Alder Wright, and Mr R Wright, who have kindly perused and corrected some of the more important sections. When it is borne in mind that the article on Acomte Bases occupies 44 pages, that on Atiopine and its Allios 27, Coca Alkaloids 23, Opium Alkaloids 67, Cinchona Alkaloids 79, and Tea and Coffee 27 pages each, it is evident that these gentlemen had no light task

I have also to acknowledge the zealous assistance of Mr G E Scott Smith, Mr C M Cames, Mr G S A Cames, and other workers in my laboratory, in researches on the Assay of Acouste Bases, the Determination of Caffeine, and much similar original experimental work, the results of which will be found duly recorded

In the sections on Tea, Copper, and Cocoa, which conclude the Volume and together occupy 73 pages, I have moorporated nearly every item of trustworthy information of a chemical nature within my knowledge, and I believe these articles will be found of service by many besides professional chemists

PART III., completing the work, will be published as soon as possible, and will, I hope, be followed at no distant date by a New Edition of the earlier Volumes.

ALFRED II. ALLEN

101, LEADENHALL STREET, LONDON, E C, 1st October 1892

CONTENTS.

AMINES AND AMMONIUM BASES	PAGE
Classification and Nomenclature of Amines,	PAGE
MONAMINES, Distinction and Separation of Monamines, 4, Methylamine, 9, Dimethylamine, 12, Trimethylamine, 12, Ethylamines, 17	3
Ammonium Bases,	18
HYDRAZINES	
Hydrazine,	22
SUBSTITUTED HYDRAZINES, Ethyl-hydrazine, 26, Phenyl-hydrazine, 27, Hydrazones, 30,	25
Osazones, 30, Pyrazolones, 30, Antipyrine, 32	
BASES FROM TAR.	
Classification of Tar Bases,	39
Aniline and its Allies,	40
Amilne, 43, Amilne-sellphome Aonis, 19, Nitranilnes, 50, Tolimidnes, 61, Xylathes, 57, Camdines, 59, Amilne Ols, 60, Amildes, 67, Acetamilde, 68, Benzamilde, 72, Substitated Amilnes, 72, Dumelhyl-amine, 74, Dipheryl- amine, 79, Amido-phenols, 80, Phenacetms, 81, Pheny- lem-diamines, 86, Escaláne, 88	

CONTENTS.

V1

Y	PAGE
NAPHTHYLAMINES AND THEIR ALLIES, a-Naphthylamine, 91, g-Naphthylamine, 92, Naphthylamines, 93, Amidonaphthols, 94	or
Pyridine Bases,	96
Pyridine, 99, Piperidine, 106; Homologues of Pyridine, 107, Pyridine-carbovylic Acids, 110, Pyriol, 113, Iodol, 111	
Quinoline and its Allies,	113
Quinoline, 116, Antipyretics allied to Quinoline, 110, Thalline, 120, Quinazolines, 122	
ACRIDINE AND TES ALLIES,	123
Acridine, 123, Phenauthridine, 126.	
,	
VEGETABLE ALKALOIDS	
CHARACTERS AND CLASSIFICATION OF ALRALOIDS,	1.27
GENERAL REACTIONS OF ALKALOIDS,	130
Reactions of the Alkalonds with Acids, 130; Thintion of Alkalonds, 130; Reactions of the Alkalonds with Alkalos, 132, Seponification of Alkalonds, 138. General Pruppi- tants of Alkalonds, 134, Colomi-teactions of Alkalonds, 144, Physiological Tests for Alkalonds, 149	
Isolation and Publification of Alkaloids,	151
Extraction by Immiscible Solvents, 154, Diagendorff's Method of Separating Alkaloids, 159	
Constitution and Synthesis of Alkaloids,	163
VOLATILE BASES OF VEGETABLE ORIGIN,	170
Conne, 171, Assay of Hemlock, 176; Lupine Alkaloids, 178, Nicotine, 179, Tobacco, 184; Snuff, 193, Priurine, 191, Lobeline, 196, Spartene, 197, Spigeline, 198.	
Acontre Bases,	198
Constitution and Characters of the Acontre Bases, 201, Aconitine, 207; Anhydro-aconitine, 213; Aconine, 214; Amorphous Bases, 215; Pscudaconitine, 216, Ventric Acid	

CONTENTS VII

PAGE

293

391

218, Japacomiane, 220, Premeoniane, 221, Lyacomiane, 223, Acolyctne, 224; Myoctonine, 225, Atisine, 226, Assay of Acomite and its Preparations, 228, Toxicology of Acomite, 236	
E AND ITS ALLIES TROPEINES,	243
statution of Atropine and its Allies, 244, Atropine, 247, Hyoszyamine, 249, Hyoszine, 250, Atropamine, 251, Belladoninie, 252, Homatropine, 253, Detection and Determination of Tropenics, 254, Belladonia, Henbane, and Stramonium, 262	
LEALOIDS,	270
ame, 273, Benzoyl-ecgonine, 282, Ecgonine, 283; Base- allied to Cocaine, 284, Amorphous Bases of Coca, 287, Coca Leaves, 290	

ATROPES

COOM A:

Coc

OPIUM ALKALOIDS,

Bases, 453

Con

Constitution of Opium Bases, 294, General Characters, 300, Coloux-reactions, 303, Separation, 305, Morphine, 304, Apouncephine, 319, Base Associates of Morphine, 304, Codeme, 321, Cryptopine, 344, Narcenine, 337, Rhosefine, 331, Theusen, 331, Opius, 331, Theusen, 331, Opius, 331, Morphine, 340, Morphine, 340, Morphine, 340, Adulterations of Opium, 340, Assay of Opium for Morphine, 342, Tincture of Opium, 360; Compound Timeture of Camphon, 363;

Toxoology of Opum and Morphine, 355

STRYCHEOS ALEADOIDS,

Strychnine, 361, Detection of Strychnine, 384, Toxicology of
Strychnine, 372, Easton's Syrup, 376, Veranu-Aillers,
378, Barena, 381, Nat. Vomane, 384, Currae, 387

CINCHORNA ALKALOIDS,
Table of Cinchona Bases, 392, General Properties of Cinchona Bases, 394; Quinnie, 397, Quinnie Shiphata, 406,
Evamination of Quinnie Salts, 408, Chrate of Iron
and Quinnie, 418; Hydioquinnie, 428, Quinniène, 429;
Quinnamie, 427, Cinchonidine, 428, Cinchonie, 431,
Amorphous Cinchona Bases, 433; Alkadodo of Remijus
Barks, 436, Cupreine, 438, Cinchona Barks, 440;
Assay of Cinchona Barks, 449, Separathon of Cinchona

Berberine and its Associates, Beibeime, 461, Ovyacanthine, 465, Hydrastine, 467, Calumba, Root, 471, Columbin, 472	PAGE 461
Cappuise and Des Allins, Chiffeun, et 44; Isolation and Determination of Califeune, 484, Theobromine, 489; Duiretin, 497, Tea, 499, Extract and Infusion of Tea, 560, Adulterations of Toa, 560, Ash of Toa, 510, Tamin in Tea, 515, Exhausted Leaves in Tea, 513, Feames of Tea, 521, Recognition of Foreign Leaves in Tea, 562, Paraguay Tea, 526, Collec, 527, Rossting of Colles, 539, Pactitious Collec, 535, (Photory, 538, Adulterations of Colles, 539, Collec Extract, 555, Kela Nuts, 554, Guarana, 555, Cocca und Chiccolite, 555, Cocca Nile, 557, Commercial Cocu, 561, Excite of Cocca, 562, Analysis of Cocca, 564, Calso Butter, 568	479
PLATES,	572
INDEX,	577
ADDENDA (sessed with the reprint of 1889),	589

AMINES AND AMMONIUM BASES.

WURTZ, in 1848, pointed out that one of the hydrogen atoms of ammonia, HaN, could be replaced by ethyl, CaHs, and shortly afterwards A W H of man n proved that the substitution by ethyl and other alkyl radicals could be extended to the second and third atoms of hydrogen, the new bodies thus produced being powerfully alkaline and in other respects closely resembling ammonia itself Hofmann called these new bases amines, and proved them to be the simplest members of a numerous class of synthetically producible compounds. He classified them as primary, secondary, and tertiary amines, according as one, two, or all three of the hydrogen atoms of the ammoniamolecule were replaced by alcoholic or alkyl radicals. As these atoms of hydrogen may be, and very often are, replaced by two or more different organic radicals, mixed amines exist, and are capable of numerous metamene modifications Thus a base having the empirical formula C5H13N may have any one of the five following constitutions -

	1	Amyl-amine,			C ₅ H ₁₁ H H
	2.	Butyl-methyl-amine, .			$\left. \begin{smallmatrix} \mathrm{C_4H_9} \\ \mathrm{CH_8} \\ \mathrm{H} \end{smallmatrix} \right\} \mathrm{N}$
	3	Propyl-ethyl-amine, .			$\begin{pmatrix} C_5H_7 \\ C_2H_5 \\ H \end{pmatrix}$ N
	4	Propyl-dimethyl-amine,			$\begin{pmatrix} C_3H_7 \\ CH_3 \\ CH_3 \end{pmatrix}N$
	5	Diethyl-methyl-amine,			$\left. \begin{array}{c} C_2H_5 \\ C_2H_5 \\ CH_8 \end{array} \right\}N$
,	ш	PART II.			

Of these metamore bases, the first only is a primary momanine, the second and third are secondary ammes, and the fourth and fifth tertanry bases. They could be distinguished by their behavious with ethyl nodule, introva and, and the other reactions described on page 4 et sec.

The hydrogen of armona may also be replaced by an actile radical, such as a cotyl or bcn.coyl, when the resultant conpound no longer possesses bear properties, and is termed an am dc (eg, acctande, $C_2\Pi_2ONH_2$) Mixed compounds also exist such as

which may be called either methyl-acetamide or acetylmethylamine. Bases are also known which are derived from the replacement of certain of the atoms of hydrogen in two, three, and even four associated molecules of ammonis, the products heinig called respectively diamino, three mines, and to train in a, which closely resemble the monumines in their general characters. The following are aximalised such lasses —

no torio utub min ordin	Pres or Sucar pube	,
Monamines Phenglamine (Anthine) (C ₆ H ₅) H H N	$\left. egin{array}{l} C_2H_5 \\ C_2H_5 \\ C_2H_5 \\ H \end{array} \right\} N$. CH_{η} CH_{η} CH_{η} CH_{η} CH_{η}
DIAMINES-		
Phenylene-diamine	Diethylene diamine	Tricthylene diamine
H,)	II _a)	(C,II,)")
$\left(C_{0}^{H_{2}}\right)^{\prime\prime}$ $\left\{N_{2}\right\}$	$\left(\begin{array}{c} \left(C_{2}H_{4}\right)^{n}\\ \left(C_{3}H_{4}\right)^{n}\end{array}\right\}N_{2}$	$\begin{pmatrix} (C_3\Pi_4)'' \\ (C_2\Pi_4)'' \\ (C_n\Pi_4)'' \end{pmatrix} N_2$
Π_2	(C ₂ H ₄)")	(C,II,)")
TRIAMINES-		
Disthylene tru	mung.	Triothylene triamine.
$\left(\begin{array}{c} \mathbf{H_5} \\ \left(\mathbf{C_2} \mathbf{H_4} \right)^n \\ \left(\mathbf{C_2} \mathbf{H_4} \right)^n \end{array} \right\}$		$(C_{3}H_{1})_{,,}$ $(C_{3}H_{1})_{,,}$ $(C_{3}H_{1})_{,,}$ $(C_{3}H_{1})_{,,}$ $(C_{3}H_{1})_{,,}$
TETRAMINES-		(0011)
	Triethylene-tetramin	a.
	$\left(C_{9}H_{4}\right)_{8}^{"}$ N_{4}	

It is evident that the formule in the text do not exhaust all possible modifications of the bese C₆H₁₅N, as they do not take into account the various isomeric modifications of which propyl, butyl, and anyl are susceptible.

Interesting bases are also obtainable by the substitution of organic radicals for the hydrogen atoms of H₃P, H₂As, and H₂Sb

The majority of the known bodies of the amine class are synthetical compounds of great scientifie but hitle practical influences. Some few ammes have been found to exist naturally in plants (\$g_1\$ times by/damne, counte), and others are met with in animal fluids (\$g_1\$ urea), or the products of the decomposition of animal matters (leatene, glycomie). The ten-bases may be regarded as a belonging to the amine class, aniline and toluid in a being primary, and pyridine and quinoline terisary monamines. Pleitidine, connine, and servocine are examples of secondary monamines, while ures and diamidoben zene may be regarded as diamnes, and bitures that guant dine as triamnes. Choline and neurine as related to the fetra-alkyl-ammonum bases. The menumines may be advantageously considered at the present stage, but the majority of the amine bases will be more conveniently desembed in other chapters.

MONAMINES.

Those bases are derived from one molecule of ammonis by the substitution of one or more of the hydrogen atoms by an equivalent number of alkyl radicals. The first body obtained of this class was et hy 1 arm in e, $C_{\rm H}$, $N_{\rm H}$, prepared by W ur v z in 1848 by distilling ethyl eynnuste with constit potesh. Me th y1 cm z1 in 1849 by distilling only expansion of the class obtained by the same chemist in the following year, by the distillation of methyl isocyanic (see fointfule) with caustic alkah — 2NCO H + CH, NCO + CM, + NH,

Hofmanı obtamed the monammes by the reaction of an althy is outdoe on an also hole solution of ammonia The seasons in not a simple one, all three monammes being formed together with a tetra-slightated ammonium base Thus, when ethyl solutio is heated with also hole ammonia to 100° in a scaled tube, there are obtained—

```
 \begin{aligned} & \text{Hydhodule of ammonia}, & & \text{H,N}, \text{H.I.} = \text{H,N} \\ & &, & \text{nonoethylamine}, & & (\text{C}_2\text{H}_2)\text{H}_3, \text{N,II} = (\text{C}_2\text{H}_2)\text{H}_3, \text{N,II} \\ &, & \text{dethylamine}, & (\text{C}_2\text{H}_2)\text{H,N}, \text{H.I.} = (\text{C}_2\text{H}_2)\text{H}_2, \text{N,II} \\ &, & \text{trethylamine}, & (\text{C}_4\text{H}_2), \text{N,II} = (\text{C}_4\text{H}_2), \text{H,I.} \\ & \text{Idode of totar-ethyl-ammonium}, & (\text{C}_4\text{H}_2), \text{N,C}, \text{I,I.} = (\text{C}_4\text{H}_2), \text{N,II} \\ & \text{Idode of totar-ethyl-ammonium}, & (\text{C}_4\text{H}_2), \text{N,C}, \text{I,I.} = (\text{C}_4\text{H}_2), \text{N,II} \\ \end{aligned}
```

Similar products result when bromide or chloride of ethyl as substituted for the iodide, except as to the relative proportions of the animes obtained. Thus chloride of ethyl produces almost exclusively EtH_NCl, with small quantities of EthH_NCl and

Et_hNG1, ethyl bromade gives chiefly EHF_hNE, with very appreciable quantities of Et_hI₂NE and Et_hINE), but very little Et_hNEr; while ethyl nodde produces EtH₂NI, Et_hI₄NI, and Et_hIIN1 in about equal proportions, as well as vary appreciable quantities of Et_hNI (Groves, Juno Chem. Soc., Mil 32).

A similar series of products is obtained by heating rodule, beromale, or intrate of methyl with a solution of aminoma in methyl alcohol. When the methyl intrate and aminoma solution are used in equivalent preportions for the nexthern—MNN+H, H,N=McH,N,HNO, monomethylamic is the chief preduct, though more or less of each of the more highly substituted, the intrata of tetra in ethyl-nam on 11 um, Mc,N,NO, is product in large excess, and the same quaternay compound is founced if methyl high methyl high substituted for the intrata of the control of the control

The complex nature of the products obtained by testing alkyl coldies, &c. with alcohole ammonia is due to the tendency of the ammes first produced to react on the renaming portions of the alkyl loddle or other salt to foun ammonium include and more highly substituted ammes. Thus—

$$\begin{array}{l} \Pi_{0}N + C_{2}\Pi_{5}I = (C_{2}\Pi_{5})H_{3}NI\\ (C_{2}H_{5})H_{2}N + C_{2}H_{6}I = (C_{2}H_{6})H_{2}NI\\ (C_{2}H_{5})_{2}IIN + C_{2}H_{5}I = (C_{0}H_{5})_{3}IINI\\ (C_{2}H_{5})_{3}N + C_{2}H_{5}I = (C_{2}\Pi_{5})_{3}NI \end{array}$$

The hydriodides of the amines similarly react with alkyl foddles in presence of ammonia to form aminonium todide and more highly substituted amines.

From these reactions it follows that the hydrodule of diethylamme, for instance, may be obtuined by heating the hounds or nodule of ethyl with a calculated amount of mono-chipanine in a sealed tube. A great variety of mixed amines may be obtained by precessly smular means

DISTINCTION AND SEPARATION OF PRIMARY, SECONDARY, AND TERTIABY MONAMINES

a It an amme be heated to 100°, under pressure, with an everest of alkyl lodded, a quaternary 10d1de will at length be formed, and the problem whether the original laws was a primary, secondary, or tertary amme will be solved by comparing the composition of the ultimate product with that of the original base or its hydrodude. Thus, if methyl nodule has been the alkyhang agent employed, the nodde of the compound ammonium ultimately obtained will differ from the hydrodude of the original

base by $3CH_{2}$ if the amine was primary, by $2CH_{2}$ if secondary, and by CH_{n} , if tertiary

b The following is an outline of the method devised by A. W Hofmann for the separation of the mixed amines resulting from heating ethyl jodide with alcoholic ammonia -The product of the reaction is filtered from ammonium iodide, which is nearly insoluble in the alcoholic liquid, and is evaporated to dryness to get rid of excess of alcohol, free ammonia, and unchanged alkyl nodide. The residue is then distilled with caustic notash, when the hydrodides of the amines are decomposed, the bases volatilising, while the indule of the tetra-alkylated ammonium base remains in the retort unchanged by, and insoluble in, the strong notash solution. The mixture of amines is conducted over caustic lime, and then condensed by passage through a well-cooled tube. The bases are then treated in a flask with one and half times their weight of ethyl oxalate (previously dried over calcium chloride), which is added gradually through a tapped funnel. This has no action on triethylamme or other tertially bases, but converts disthylamine into hound ethyl diethyl-oxamate, and mono-othylamine into solid diethyl-oxamide, according to the following equations .-

1.
$$(C_0H_0)_0C_0O_4 + 2(C_2H_0)NH_0 = C_2O_3\left\{\frac{NH(C_0H_0)}{NH(C_2H_0)} + 2(C_2H_0)OH\right\}$$
Eithyl analata. Biblyminno Diethyl counside Alcohol

2. $(C_2H_0)_2C_2O_4 + (C_2H_0)_2NH = C_2O_3\left\{\frac{OC_2H_0}{N(C_2H_0)_2} + (C_2H_0)OH\right\}$
Eithyl consiste Dictabilantine Eithyl Dictabilation annalate Alcohol

The inpud gets very bot, but for the completion of the reaction the mixture should be heated to 100° for several days in a closed vessel. The trichtylamine, which has taken no part in the reaction, is then distilled off on the water-bath. The iscalar is well cooled, and the sold oxamide separated from the hugud oxamide by pressure? On subsequent distillation with caustice potach,

¹ Diethyloxamale may also be separated from the othyl diethyloxamale by coll water, in which the former dissolves oxely, the latter very spaningly. If how water be used, the separation as more perfect and the readual oximate quite pure, but some of it suffers hydrolysis and goes into solution as distriploxamic acade.

 2 Some ethyl monosthyloxamate, $C_2O_2\left\{ \begin{array}{ll} O.C_1H_8\\NH.C_2H_8 \end{array} \right.$, is always formed from the primary animes in this reaction

these compounds yield the primary and secondary assures respectively ----

1, C₂O₄(NH C₂H₆)₂+2H (OK)=C₂O₄(OK)₂+2H(NH C₂H₄)

2.
$$C_2O_2$$
 $\left\{ \begin{array}{l} O(C_2H_o) \\ N(C,H) \end{array} \right\} + 2H(OK) = C_2O_2(OK)_2 + H.N(C_2H_o)_2 + HO(C_2H_o)$

The foregoing process is available, with critain modifications in detail, for the separation of the annies of methyl and other homelogues of chtyl, and, in fact, is of general application for the separation of primary, secondary, and teatrary annies, the first clusforming examides, the second examic ethers, and the third being unacted on by chilfy examine.

An important modification in the foregoing method has been made by Duvillies and Bussine (Ann. Chim Phys. [5], xxiii. 289), who operate on an aqueous solution of the bases. Under these conditions, the minuary amines are converted by ethyl oxalite mto insoluble or sparingly soluble ovanides, while the secondary and tertiary bases are unchanged, or at any rate remain wholly in solution. After separating the oxamides by hitration, the motherhonor I is boiled for some time, which causes the hydrolysis of the ethyl diethyloxamate with formation of diethyloxamic acid, (CoH.) N CoO OH, and the further change of this into the acid oxalate of diethylamine, (Calla), HN H, CaO, 1 This salt separates on cooling, and yields the free base on distillation with alkalı The filtrate is distilled with potash, the loses dried by caustic potash, and dissolved in absolute alcohol. On adding ethyl exalate to this solution the secondary amines are converted into examic others, while any remaining primary animes are converted into the corresponding examides. After allowing the inixture to stand for twenty-four hours to complete the reaction, the alcohol and unchanged tertiany bases are distilled off on the water-bath. The examates remaining in the retort may be converted into calcium salts by treatment with milk of lime or the secondary bases at once liberated and recovered by distribution with caustic potash.2

¹ The treatment described in the brackets is optional, and chirfly of advantage in the separation of ethylamines.

⁵ The convenion into addition sails is especially satisful for the trustment of the citylinens. The progratistic alexam divelylenamels and memority-carnate are filtered off, and the lithasis heeted still alcohol, which prespective the remainder of the column sails. The precipitative are treated with boiling water, when the monosthylaximate desawlers, and is deposited again on cooling in large crystals, which on distillation with potenty light displaname. On concentrating and cooling the models—ilgues, calcium distillations is paratise. It is recrystallised from alcohol, washed with other to nanove my calcium and coloning and distilled with potch when it publis pare diskiplaname, adhering oxamide, and distilled with potch with my though pare diskiplaname.

Duviller and Dusane have applied this method to the analysis of the complex mixture of amines present in commercial timethylamine from vinasses (page 13) A Muller (Bull Soc. Ohim, kin 202; Jour Ohem Soc., kivin 501) has described a method for the separation of amines based on much the same principle

The primary, secondary, and testing monamines may also be distinguished by the following reactions —

- c If a primary monamine be boiled with alcoholic potash and chloreform, the characteristic and highly disagreeable odour of the corresponding car be mine or is onitrile is evolved, according to the reaction —MeNH_e+CHCl₄+8KHO = MeNC+3H₄O+3KCl.
- d. If a primary fatty monamine be dissolved in a mixture of equal measures of alcohol and carbon dissiphate, and the liquid thus boiled down to one-half, at hio carbo mate will be formed thus $2\text{MeNH}_2 + \text{CS}_2 = \text{MeNH CS S NMeH}_3$

If the resultant liquid be boiled with a solution of mercuric or ferric chloride, a pungent odom of mustard oil will be produced. owing to the formation of an alkyl iso-thiocyanate 1.—

MeNH CS S NMeH, + HgCl, = HgS + MeNCS + NMeH, Cl

e Nitrous seid converts pumen y fatty monammes unto the corresponding a l c a ho is — MeH_N+NO OH = Ma OH+OH_+N_3 Atomatic primary ammes (eg, amline) are converted by nitrous and into diaso-compounds —PhNH_2+NO OH=PhN NOH+H_2O

Secondary amines, whether fatty or anomatic, are converted by mitroue acid mit on it re s an in es, thus — Ma,NH + NO OH = Me,N NO+ H₂O The mitrosemines are yellow inquids, of neutral character and aromatic colour, volatile without decomposition at current of stoam Weak reducing agents convert them into hydrasines (page 27); but by more powerful hydrogening agents, or by warruing with alcohol and hydrochloric acid, they are reconverted into the original secondary amines

Nitrous and has no action on tertiany fatty amines. It converts most tentiary aromatic amines into nitroso-derivatives which still possess basic properties

- In practice, the action of introns and on the amines is best effected by distilling their hydrochlorides with a strong solution of poissaum or sodium intrite. If a mixture of the hydrochlorides of the three methylamines be thus treated, the monomethylamines as destroyed (with formation of methyl alcohol, which will be found
 - In the case of aromatic primary amines, the product is usually a thic-treat, which requires to be treated with phosphone pentoxide to obtain the iso-thic cyanate.

in the distillate), dimethylamine is conveited into dimethylnitrosamine, which distils, while the hybrochloride of trimethylamine remains in the retort (mixed with excess of the metallic mitrie), and on distilling it with causic alkali the free base son he obtained

f Both primary and secondary monantnes react with ablehydes to form indifferent bodies. The reaction between consulted and more and dispatch planning respectively is as follows.

$$H_2 N CH_3 + C_6H_{15} CHO = H_2O + C_6II_{12} CH NCII_3$$
, and
 $2HN(CH_3)_1 + C_2H_{12} CHO = H_0O + C_2II_{12} CHIN(CH_3)_1$

This reaction has been utilised by Schriff (Aimalor, ch., 168) for the volumetre assay of numes. The base is disabled for the rolumenter assay of numes The base is disabled and the national constitution in bearage deposed in from a bantet as long as with continues to separate. Each addition of the camental solution produces a turbular from separation of satic, but this is absorbed by the calcium chloride on gentle agistism. As a primary ammerisativity from separation of satic, but this is absorbed by the calcium chloride on gentle agistism. As a primary combinate with the control of t

g The actd ferroeyanides of the tertiony antibles are remarkably usoluble in water. They are precipitated on adding potassum ferroeyanide to the solutions of the amines a udulated with hydrochloric acid. This bases can be recovered from their formsynamides by teating the precipitate with solution of cupits, shiphate, filtering, and removing the sulphuric acid and excess of copper from the filtsate by harris-water.

GENERIC CHARACTERS OF MONAMINES.

The monatonics, as a class, are resultly volatile inquids, of lower specific gravity than water. Their boiling-points has with the number of cartion atoms in the molecule. They are inflammable, burning with a yellow fiame, and the lower members dis-cive with great facility in water, forming strongly alkaline higuids of an ammoniacal order. From their solutions, ethylamine and the higher homologues can be separated by attenting the inquid with caustic potash. By boiling the aqueous solutions of the lice

1 On separating the miscosmume, which forms a yellow oil, from the aqueous distillate, treating it with aqueous hydrochlonic acid, and then reasing hydrochlonic acid gas till the liquid is homogeneous, the hydrochlonic of the secondary amme is formed, and may be obtained by evaporation of the solution.

bases, or of their salts after adding excess of lime or fixed easily alkul, the monamines can be completely volatilised, and condensed again in water or acid, and triated in the same manner as ammonia. The monamines are all powerful bases, closely resembling ammonia in their general channeters. They form crystallisable salts, and yield obloroplatinates, oblor-crystallisable salts, and yield obloroplatinates, oblor-missibilition to the corresponding compounds of ammonia. The monamies prespitate magnesium salts, but the precipitated magnesium salts, but the precipitated magnesium salts, but the precipitated magnesium hydroxide dissolves in the amine hydroxide, forming a double salt from the solution of which phesphate of sodium precipitates an amino-magnesium phosphate. The amines thus behave exactly in the same manner as ammonia.

The only ammes (not described in other chaptens) requiring detailed consideration are the primary, secondary, and tetracy monamies of methyl and ethyl These bodies are typical of the names generally, and most of the statements made respecting them would be true of all the botics of the class. Their aqueous solutions dissolve silver chlouds, and behave in much the same manner as aminoma with metallic salts, but there are some interesting differences, as shown in the table on next page, from which it will be seen that certain of the prequistates which are soluble in excess of animonia are undussolved by the annues, and vice series.

In all eases a solution of aluminium phosphate in hydrochloric acid behaves similarly to a solution of aluminium chloride (Taylor)

Methylamine. Monomethylamine

$$CH_sN = H_s$$
 H
 II

Methylamine exists leady-formed in Mercuralts assua and M personns, and so obtained (in an impure state) from these pluits, was foinedly known as mercurialine. It also exists in herring-linine, could-tu, hone-of, and the products of the distillation of wood,² bectious molesses (wasses), and cattain alkalouls

¹ The author is indebted to Leo Taylor for repeating and enlarging on the experiments of Vincont, on whose observations the table is chiefly feathful. Several blanks in the observations of Vincont have been filled by Taylor.

² The presence of the amines of methyl in pyroligneous acid and wood spirit is probably due to the reaction of actions and aminomia $-C_2H_0O + NH_0 - C_2H_0 + C_2H_0 + NH_0 - C_2H_0 + C_2H$

Metallie Salt	Ammonta H ₃ N	$\begin{array}{c} Bthylamme\\ (C_2H_4)H_2N \end{array}$	Metholamone (CH ₂)U_N	Demethyl- gustus (CH ₂) ₂ UN	I'r imethyl- amine (CH ₄) ₂ N
Aluminium	Insoluble (nearly)	Soluble	Soluble	Soluble	Soluble
Cohalt	Blue precrpi- tato, sol- uble in ex cess to brown solution.	Insoluble	Blue, insoluble in oxoses, tauned lnownish on heating	filue, meel uble in excess, tmned brownish on heating	Blue, insol- uble in excess; funced brownish on heating
Nickel	Seluble in evers to * violet-blue selution	Inseluble.	Apple green, insoluble ru oxpess	Apple meens insoluble in excess	Apple green insoluble in excuss
Zinc.	Very soluble	Selmble.	Soluble in large ex- cess, re- ppted on heating	Soluble in 1 ugo ev coss, 10 pated on heating	Soluble in very large excess, repoted on heating
Cadmitum	Soluble	Inscluble	Insoluble	Insoluble,	Insoluble
Salver	Brownish; very sol- unle in ex- coss	-	Brownish, soluble in large ex cess, 10 pited on warning	Brownish, soluble in large ex- cess, re- paired on warming.	Dity brown into changing to black, soil large everas to dark soin tion, re- pyted on warming
Cuprio,	Bine , sol- uble in everse to deep bine solution	Solublo with difficulty in excess	Bine; soluble In large our se to deep blue solution; reputed dirty brown on builing	Blue, partly soluble in ingo ex- cuss, re- puted dirty hiown on builing	Blue, partly soluble in large ex- cess, re- ppt d dirt brown on boiling
Meacurie	White	-	White , in soluble	White, sol uble in much water	Yellow, changing to very pale yel- lew
Stannie	Inseluble	Very soluble in excess		Boluble.	Solulilo.
Autimonic				Soluble	Soluble in large ex- eess
Gold	Insolubia	Soluble.	Brownish yellow ppt, tondily sol uplo lu excess to orange-red liquid	exces to brown	
Butherium,	Insoluble	Solublo			
Lend	Insoluble		Insoluble	Insoluble	Insoluble

(e.g., morphine, codene) It is also produced when caffeino is boiled with baryta-water, and by heating hydrochloride of trimethylamine to 2887, when methyl chloride and trimethylamine volatiles, and methylamine hydrochloride (mixed with some ammonium chloride) iemains

Methylamme may be prepared by the action of alcoholuc ammons on methyl lodids, but in this case dimethylamne and immethylamne are also produced (rage 3), and the man product is solded of tetramethyl-ammonium. Methylamnes is beobtained pure by treating one equivalent of acctamide with two equivalents of bromme, and then adding a 10 per cent solution of caustic potash till the colour of the bromine has nearly disampeared.—

 $\label{eq:c2H2ONH2} {\rm C_2H_3O~N\,H_2 + Br_2 + 2KHO = C_2H_3O~N~BrK + KBr + 2H_2O}\,.$

Three additional equivalents of caustic poissit are now dissolved to a 10 per cart solution, and heated in a rector to 70° C. The product of the first reaction is then gradually added through the tubulure. The gases evolved are collected in hydrochloric early and on evaporating the solution a mixture of the hydrochlorides of amnonia and methylamine is obtained, from which the latter only is dissolved by absolute abolich Of distillation with caustic alkali or slaked lime the salt yields the base, quite free from the or termedial anne.

Metilylamine boils only a few degrees above zero, and hence is a gas at ordininy temperature. One roliume of water at 12° C. dissolves 1150 measures of the gas, and hence it is more soluble even than amount, which metilylamine closely resembles in colour and general characters, but is distinguished by its ready indiamount of the content of the conte

On passing a succession of electric sparks through unsethylanium, by dx or y and z of x me thy z and z as y produced, and this is decomposed by a continuation of the treatment, with formation of a tarry deposit. When passed through a red-hot table, methyland is decomposed with foundation of hydrogen and a mmonium z and z are z

The behaviour of methylamine with metallic solutions (page 10) and various other of its reactions have already been described. It forms a series of readily crystallisable salts. The chloro-

1 The reaction which occurs is very complex (A W Hofmann, Ber., vv. 765), but the main decomposition may be expressed as follows —

CH3 CO NKBr+2HOK -CO(OK)3+KBr+CH3 NH2.

platinate, (MeH₃N)₂PtCl₁₀, is insoluble in alcohol, but soluble in boiling water, crystallising on cooling in beautiful golden-yellow scales

A method for the proximate analysis of the bases present in crude methylamine, based on the principles of the process described on page 6, has been described by A Mullei (*Bull. Soc. Chim*, xh. 202, *Journ Chem Soc*, xlvni. 501).

Dimethylamine.

$$C_2H_7N = CII_3$$
 CII_3
 N

Dimethylamine occurs in Penavian guano and pyrolignous acid, and is also present in the products of the distillation of unasses.

Dimethylamme is readily separated from the permany and tort up to methylamme by converting it une of the pld time is the plant in a to (pages 5, 6, 14), or into difficult with other plant in the plant in (page 7), on absulting the first of those derivatives with careful entering the second with fruming hydrodilous and, the dimethylamme is regionated. The base may also be obtained pure by boiling 35 parts of interso-dimethylamiline hydrodilouside with a solution of 15 hards of careful orders in order that the solution of 15 hards of careful orders in order that is solution of 15 hards of careful orders in the first plant in the first plant in the plant is the solution of 15 hards of careful orders in the first plant in the first plant in the plant is the plant in the plant in the plant in the plant is the plant in the plant in

 $C_0H_4(NO) NMe_{2}HCl+KHO = KCl+C_0H_4(NO)OH+HNMe_{2}$

Dimothylamino boils at 8°-9° C, and closely resembles the primary and tentany modelylamines. From this former it is at once distinguished by the new formation of a pieceptatic on the addition of ethyl oxalate to the autoous solution of the base (lago 6), and the non-production of an isomitale on teachmost with alcoholo potash and chloroform. From termethylamino it is distinguished by the formation of a nitrosamino on treating it with intions said, or one of its salts with a intrice (page 7).

The chloroplatinate, (Mc2II2N)2PtCl6, crystallises in very long needles.

Trimethylamine.

$$C_8H_9N = CH_9 \atop CII_1$$
 N.

Trimethylamina, often impropedly called propylamina, a base having the constitution $(C_2H_2)\Pi$ N, occurs somewhat frequently both in the animal and vegetable kingdom. In the former it occurs notably in herring-brine, and has been detected in mine, imputio-

fied blood of the calf; cod-lives oil, and other animal fluids. In the vegetable kingdom, trinctlylamine occurs in the Chenopodium valuavia (striking goose-foot), from the leaves of which it constantly exules, Arnaca montana, Macarraine anima, the blossons of the pear, white-them (Gatagine sogicandia), hawthern, and wild cherry, and in ergot¹ and other parasites of the vegetable kingdom. Trimethylamine is also a product of the dry distillation of certain alkaloids, wood, &c, but especially of the unasses or residue left after the distillation of the spirit from formented bectroot molesses. The bases obtained by the destructive distillation of this product are derived from the bet in no Q.d.J.m.NO; contained in the molesses, and conest chiefly of the menamines of methyl, among which trimethylamine predominates ²

The products of the destructive distillation of the "unasses," left after the distillation of the fermantial bestoot-molasses, vary with the concentration of the liquid As the proportion of water decreases, the quantity of ammons moreoses, and the tunnethylamine is replaced by the primary and secondary methylamines. The winasses from different localities yield varying proportions of gaseous and liquid products on distillation, the nutriles and methylic alcohol appearing to be the most variable constituents, if

 1 The trimethylamino of eigot is probably a decomposition-product of choline, $(CH_5)_5 N(C_5 H_4 O H)$ OH.

² The vinasses, or spent wash from the stills, is evaporated till it acquires a specific gravity of 1'31, when it is subjected to day distillation in cast-iron retorts The aqueous portion of the distillate contains - Ammonium car bonate, sulphydrate and cyanide, mothyl alcohol, methyl eulphido, and methyl evanide, various other bodies of the fatty series, and a large proportion of salts of trimethylamine The tar yields, on distillation -ammoniacal liquor. various oils, pyridine bases, solid hydrocarbons, phenols, and pitch of superior quality. The aqueous liquid is neutralised with sulphurio acid and concentrated, when crystals of ammonium sulphate are deposited, and vapours of methyl alcohol are evolved together with methyl cyanide and other nitules The methyl cyanide is converted in ammonia and acotate by treatment with an alkalı :-- CH, NC+NaHO+H,O-II,N+CH, COONa The dark-coloured mother-liquors retain the trimethylamine sulphate, which is decomposed by distillation with lune, the vapours being passed into hydrochloric acid. The resultant solution is botled down till the temperature reaches 140° C Ammonium chloride crystallises out on cooling, and the mother-liquor is esparated and concontrated till the boiling-point rises to 200°, the product forming commercial hydrochloride of trimethylamine, from which the free base may readily be obtained by treatment with hime or caustic alkali.

In a epecimen of "commercial trimethylamine," prepared from vinasces, Duvillier and Buisine found only from 5 to 10 per cent of tunnethylamine and come 50 per cent of dimethylamine; while the remainder consistency of methylamine, propylamino, and isolutylamine in about equal proportions;

Trimethylamine has a specific gravity of 0 673 at 0°, and boils between 9° and 10° C When pure and concentrated, trangethylthe othylamino being estimated at about 2 per cent, and ammonia being absent (Compt Rend , lvxxx. 48) The method employed by these chemists for the separation of the aminos in question was as tollows (Ann Chim Phys., [5], rrm 289) .- The agreeous solution of the free bases was treated with ethyl ovalate, the dense white precipitate of oxumides filtered off, the filtrate concentrated by distillation, and the further precipitate added to that meyiously obtained By treating the precipitate with het water it was separated into three fractions The most su-cluble portion (1) consisted of dibuty! oxamide for nossibly di-sacbutyloxamido), which melted and floated on the hot water, and on cooling formed a solid wavy mass. When iconstalhaed from alcohol, it was obtained in posity needles. The butulamine. C.H.NH. oldamed by distilling the examile with potash, had a family aromatic odour, and yielded a slightly soluble chloroplatinate, crystallising in orange-coloured plates Of the examindes soluble in boiling water, the dimensil compound (2) was first deposited. It crystalled from alcohol in hearly needles molting at 110°, and the propylamine, C.H., NH, obtained from it gave an orange chloroplatinate When the proportion of butylamine and propylamine was small, the authors preferred to utilize the comparative insolubility of their sulphates in alcohol to separate them from the other annues The most soluble portion of the mixed examiles (8) was deposited in opinion white needles or grams, and consisted of dimothyloxamide. The base obtained by distribug it with potash was converted into the sulphite, which on treatment with horling absolute alcohol was obtained quite jure, and vielded nure methulamine on treatment with notash

The mother-bytes separated from the examities of the prumary unines was distilled with easter potates, and the during ges collected in absolute aisolated. A portion of the solution was then thatted with standard areal, and the remanded gradually adoled to a quantity of eithyl evaluate sufficient in the reaction—MagNII+82,C₃O₂—(MolIN),C₃O₃2+2ReOII, assuming the alkalmity, to be whichly due to dimethylamine. The operation was conducted in a first, which was sminomeded with two and continually lanken. When the extreme completed, the finds was bested on the water both, and the alcohol and unimplicated the first was bested on the water both, and the alcohol and unimputed to the continual of the continual of

The grupy reaches left in the fasks after the datilitation of the alcohol and trimedly learns constated of the ethyl finkly lated-exameter, with threes of ribly menally sketch exameter sand exameles of primary amines. It was tearted with water, whole caused hybridyeas, and, on mentralweig the hybrid with matter, whole caused hybridyeas and, and the thours down, which on datallation with portal yielded diplement, $O(l_1, Nll_2, on all propylements, Old, Nll_2, on all propylements, Old, Nll_2, on the twing the fiftenties from the calcum examets a period with an equal colume of alcohol, a precipitate was faunch from which warm water extincted as let us un time thy lex a nate, youlding datifyliname, (CHANII, on dividiation with potats, while the less soluble portion convexted of calcum memorathyly awantay politing antifyliname under samiler treatment.$

Ethylamus, which escaped detection on Duvillier and Buisine's inst

amine is stated to have a purely ammoniacal odour; but when highly diluted, the vapour has at the same time a small of ammonia and a peculiar fishy odour suggestive of herring-brine. The latter odour is gradually developed by adding lime to a solution of the base, but requires some time to reach its maximum intensity (I. Taylor)

Trimethylamine is apparently soluble in all proportions of cold water 1

A mixture of equal measures of trimcthylamine and water is inflammable.

Timethylamme is employed for preparing pure potassum carbonate from the chlorde by a method enalogous to the ammonissods process Ammonia is not available, because of the nearly equal solicitative in water of mamonium chlorde and acid potassum carbonate, whereas the hydrochloride of trimethylamine is much more solicitie.

Trunceltylamine might, psind faces, be supposed the active agent in W oll the sun's sprocess of treating sewage with herring-brine and lime (Eng Patent No 15321, 1889), but those who have investigated the matter meline to the opinion that the bactericule is a hitherto unisolated body thay term am in oil, produced by the action of lime on one of the amines of herring-brine. Pure trumshtylamine employed without lime has not the same effect

Trimethylamine is distinguished from the primary and secondary methylamines by its negative reaction with alcoholic potash and chloroform (page 7), ethyl oxalate (page 5), and nitrous acid (page 7), and by its solution in scores of hydrochloric acid being precipitated by potassium ferrocyamide (page 8)

Trimethylamine has been employed in medicine, and is said to have proved of value in the treatment of gout and acute rheumatism

examination of the bases from ynnews, owing to the small proportion present, was subsequently detected by distilling with pleased the mother-lapsors obtained by treating the oramides with water, and convexing the bases insulptate. On treating these own subplicate contracting these with absolute alcohol, the subplicate of methyl-amino remained. On distilling the solubile portion with alkali, collecting the bases in takeolite alcohol, and treating the solution with child collation, where the subsect of the subsec

According to Guthii, the isolubility of transitylamme in water is notably dummahed by heating, the liquid becoming distinctly turbid comparts nicotine) from partial separation of the base. Thus a 10 per cent solution of transitylammo in water became turbid at 22° C, as 5 per cent at 22° 5, and 4 per cent solution at about 42° C. Leo Taylor has failed to confine Guthine's observations, which were not improbably made on impure material. (A valuable description of its therapeutic effects will be found in the Year-Book of Pharmacy for 1873, pages 197-262)¹

Trunshylamane combines with outlon dissiphate at the columny temperature with great evolution of bast, ascording to the equation CS,+(CH_A)N=N(CH_A)CS S CH_B. The produce, which may be regarded as the une taby it the carba must actual prepared more resultly by passing gaseous frinchlylamine into a maxime of carbon desulphate and also he. It is obtained on acapaning the solvent in white thombie necdles, melts at 125°, and decompose gradually at the ordinary temperature. It is solvible in distasticated and water, but nearly insolution in also discontinuous control of the colored production of the colored production of the colored production of passing of the colored production of the colored production of the colored production of the colored production of passing of the colored production of the colored product

Trunchylamus Hybrochlards Hydrochlorate of trunchlyl, anno Chlordto of Trunchylamunoum (Ghyl, MINO! This salt is obtained by neutralising trunchylamune with hydrochlora card is differs from aminomum chlorde in being extremely deliquis-cents, and soluble in absolute alcohol. The fi-ly obour of the base bleated on tracting the salt with line or caustic alkali further distinguishes it from ammonium oldroids. With plature chloride it unites to form the rhibo-appliantate, (Mc, HIN, PhS), a compound which optabilizes in orange octobeths, sparnigly soluble in skenitra clachd.

When heated to 260°-285° C, trimethylamine hydrochlorule is decomposed with formation of free trimethylamine, ammonia, and methyl chlorido:—

$$3Me\Pi NCl = 2Me_aN + II_aN + 3MeCl$$
.

This reaction has been utilised by Camillo Vincent for the manineture of methyl chlorude. The vapours are passed through hydrooblone acid, which absorbs the bases, while the gracous mothyl chloride passes on. It is washed by dilute causics code and direled by strong sulphurus each, after which it is collected in a gas-blurus each, after which it is collected in a gas-blurus each after which it is condensed to lequal. The vapour of liquid methyl chloride has a tension of 2.5 atmospheres at 0 and 4.8 at 20° C.

¹ The solution of timestlylamne for medicand use should be clast, solour-less, and of 1 124 speeds gravity. It should have a position relour, recaling that of amonon and harmy-lims, he meetids in all proportions with water and alcolo, and contain 29 per cent of the base. One measure of hydrochron act, of 170 speeds gravity, should neutrinos blue measures of the solution of the base, and the salt obtained on evaporating the resultant solution should be committed without a modulet should.

Methyl chloride is extensively used in the annine-dye manufacture for preparing methylaniline and dimethylaniline, which compounds form the starting-points of numerous colouring matters.

Ethylamines.

The amnes of ethyl are obtainable in the manner already described (page 3). A convenient source of the primary amine, C₂H₁NH₂, is the crude ethyl chloride obtained as a bye-product in the manufacture of chloral (A. W H of m in , Ber, in 10-99, 776). When ethyl chloride is heated to 90° under pressure with an equivalent proportion of stong aqueous aminona, a layer of trachylamine containing ammona is formed, while the squeous liquid contains the hydrochlorides of ethylamine and ducthylamine. When a similar mixture of aqueous ammona and ethyl chiloride is heated under pressure to 160° C, H,NCI, BH,NCI, and Ex,NCI are the chief products, only traces of Et₂H₂NCI and Et₂HNCI being formed.

The ammes of sthyl can be separated by methods already described. They present the closest analogy to the corresponding methyl bases. Various differences between the three ammes are described on page 4 st seq. The following table shows other of their characteristic properties.

	ETHYLAMINE	DIETH) LAMINE	TRIETHYLAMINU
Formula,	(C ₂ H ₆)H ₂ N	(C,H ₅),HN	(C,H,hN
Boiling-point, * C	19	66	90
Specific gravity	80 6864	<u>0</u> 0 7000	20 ₀ 7277
	20 708	20 708	
Reaction with zino sulphate	Precipitate soluble in excess	Proofpitate insoluble in excess	Precipitato in-
Product when holled with nitions achi (or a salt of the bases with sodium nitrate solution)	Alcohol and mino	Diothylnitrosamine, a neutral oily liquid boiling at 177, and distilling with steam (pago 7).	Unchanged
Hydrochloride	Deliquescent lamina and prisms	Non deliquoscent plates	Non deliquescent lamina
Platinichloride	Hexagonal rhom- bohedis, mode- rately soluble in water	Monoclinic, mode- rately soluble	Monoclinie, very soluble.
Aoid ferrocyanida	Soluble.	Soluble	Very sparingly soluble

AMMONIUM BASES.

By the actors of excess of an alkyl solide on ammona or an amme, all the hydrogen atoms of ammona can be taplaced by alkyl radicals, the tertany ammest thus formed combaring with another molecule of alkyl nodulo to produce the tool do of a tetra-alkylated an unrount without methyl solide has acted on ammona, the product is tetamethyl-ammonium solide, (CH₂)₂NI, but by obvious modifications in the process, similar compounds containing other alkyl-radicals can be obtained. Thus, H of mann prepared the indicate of methyl-cityl-tamyl-phenyl-ammonium—(CHI)₂(CH₂)₂(N₂(H₂)₃(N₂(H₂))₃(N₂(H₂)₃(N₂(H₂))₃(N₂(H₂))₃(N₂(H₂))₄(H₂(H₂))₄(N₂(H₂))₄(N₂(H₂))₄(N₂(H₂))₄(N₂(H₂))₄(N₂(H₂))₄(N₂(H₂))₄(N

The same product results from the action of ethyl rollide on truncthylaume as by the action of methyl nelide on dimethylsthylamne. This fact proves that the body formed is not merely a molecular compound of the constitution.

$$\begin{array}{c} \operatorname{CH}_{q} \\ \operatorname{CH}_{1} \\ \operatorname{CH}_{3} \end{array} \} \operatorname{N}_{1} \operatorname{C}_{2} \operatorname{\Pi}_{0} \operatorname{I} \text{ , or } \begin{array}{c} \operatorname{CH}_{q} \\ \operatorname{CH}_{1} \\ \operatorname{C}_{2} \operatorname{H}_{5} \end{array} \right\} \operatorname{N}_{1} \operatorname{CH}_{1} \operatorname{I} \text{ , }$$

but that it is the true redide of a tetra-alkylated ammonium ---

The identity of these and similar compounds furnishes important evidence of the pentavalent character of introgen

The robbles of the tetra-alkylated anniousnums are quite macted on by caustic potent even in beaung, but we twill recently preceptated argentic exists to four noticle of silver and the hydroxides of the tetra-alkylated anniousnums. These hydroxides are non-volatile, sympy or solid debigo-semi substances, of highly caustic, alkaline changetes, presenting, as a class, a strong analogy to caustic potash. Many of them have marked poisonous characters.

Such of the natural vegetable alkaloids as have the constitution of terhary bases unite with alkyl nodules to form compounds which have the characters of todales of compound unmonutum, from which the corresponding hydroxides can be prepared, real-above described, by reaction with oxide of alver. Thus, for example, from more plains, $C_{\rm pl} I_{\rm pp} NO_{\rm p}$ may be prepared—oxample, from more plains, $C_{\rm pl} I_{\rm pp} NO_{\rm p}$ may be prepared—

These bodies are sometimes formulated and described as the hydrodude and hydrate of ethylmorphine, $C_{17}II_{18}(C_2H_5)NO_3$; but such a view is inconsistent with their characters

Similar bodies are obtained by action of alkyl bodies on stry of h in a. The hydroxides of methyl—and ethyl-strychnium ($C_{\rm D}H_{\rm m}{\rm MeNO}_{\rm S}{\rm OH}$ and $C_{\rm m}H_{\rm m}{\rm ENO}_{\rm S}{\rm OH}$) are stong, very soluble bases, which form earbonates and precentate metallic hydroxides from metallic solutions. In their physiological action they smulate the paralysing action of curarine rather than the tetanic poisoning of strychnium tiself

Similar bases can be obtained by the action of alkyl salts on dimmines or ammonia. Thus, as and-product of the action of excess of sthylene dibrounde on ammonia is tetra - ethylenedi-ammonium-dibrounde (C_HH_c)_cN_cB_E, from which the hydroxide, (C_HH_c)_cN_cH_c can be obtained by treatment with oxide of silver. This base is a powerful causine alkali and nonvolatile

Choline and neurine, described in the chapter on "Animal Bases," are natural products having the constitution of ammonium bases. Thus —

It will be observed that neurine and choline only differ from each other by the elements of water

Bases of similar characters and constitution have been prepared, containing phosphorus, arsenic, or autimony in place of nitrogen Thus, there have been obtained —

Tetramethyl-ammonium hydroxide, Me₂N OH
Trimethyl-ethyl-phosphonium hydroxide, Me₂EP OH
Tetrethyl-arsonium hydroxide, Et₂As OH
Tetrethyl-sthonium hydroxide, Et₃Sb OH

Tetrethyl-ammonium Compounds.

When perfectly anhydrous ethyl rodule is added to trimethylamine previously dired over caustic potash, combination gradually occurs with evolution of heat, and in a few days the mixture sets to a solid mass of

Tetrethylammonium Iodude, (C₂H₅)₄NI This compound is preferably prepared by exposing a mixture of equivalent proportions of triethylamine and ethyl rodule to a temperature of 100° for a fow muntes in a flask furnished with a well-cooled invorted condensor, or preferably in a scaled tabe. Violent reaction ensuses, and, on cooling, the product sets to a hard mass of crystals. On dissolving the mass in water, and allowing the solution to evaporate spontaneously, the todde is obtained in extensicly bitter cytals of considerable size, which, when pure, are colourless, but are apt to be mixed with reddish crystals of the tri-rodiide, (GJH₂),NII,1

Tetesthylammonum todde as not volatile at 100°C, but when apidly heated in a retort to a higher temperature it melts and suffices decomposition into ethyl i odiido and triinothylamine, which form separate layers in the receiver but re-unite to noduce the original composition.

Tetesthylammonum iodule is wholly undecomposed by treatment with cariste poteah or soda, but is much less soluble in caustic alkaline solutions than in water. Hence, on adding excess of causter potent to the concentrated aqueous solution, a solid crystalline mass is produced. This behaviour sharply distinguishes the iodide of test-stily-lammonium (and of other compound insenniums) from the compounds Eq. ITM, Eq. IIM, and ISH, NJ, which are at once decomposed by caustic alkali, with theiration of the corresponding minus. The aqueous solution of test-shipiammonium iodic vestes with agentic naturator or sulphato to form a presuptate of argentic codde and a solution of the test-shipiammonium intrate or sulphate

Therefithamonium Hydrogues, Call Jan Oll, is oblamed in solution by adding freshly-precupitated coate of silver to a dilute and warm solution of tethethylaminonium iorbide, until the become colour of the silver existe coses to change into the lemon-yellow of the solution. The solution is then filtered, and may be evaporated to a considerable setten at a gentle heat, but further concentrium must be conducted in scase, at the ordinary temperature, over sufficient to the conducted in scase, at the ordinary temperature, over sufficient to the solution of the base are deposited, but those subsequently disappear, and the hapud ultimately dries up to a seem-solul mass of

Tetrebylaumonum hydroxide presents the closest analogy to caustic potash It is inglish deliquescent, alpostis acribin duxido from the air, and the squeous solution has a strong alkalium reaction. It has an alkaline, caussic, and axtremely lutter taste, and in a concentrated state burns the tangue and ects on the skin like caustic plotash. With metallic solutions it behaves like the caustic alkaline, except that alumnium hydroxide is solution with

¹ This compound is readily obtained by dissolving reduce in a solution of tetrethylammonium reduce.

difficulty in excess of the reagent, and chromic hydroxide is quite insoluble

A moderately strong solution of teleshylammonium hydroxidate, may be boiled without decomposition, but in a concentrated act, even at 100°, the liquid fields strongly, and the base is resolved gradually but completely into title hylamine, ethylene, and water.—CLFLNO HI — (CJFL)N + CLFL + HOH!. This reaction affords a convenient means of obtaining triethylamine unmixed with the primary and secondary amost

When a solution of tetrethylammonum hydroxide is boiled with a slight excess of othyl lodule for twenty-four hours, under a reflux condenser, the solution becomes perfectly neutral, the following reaction occurring $-(C_2H_L)NOH + C_2H_L = (C_2H_L)NII + C_2H_COH$

Tetrethylammonium hydroxide also hydrolyses ethyl oxalate, and saponifies fats as readily as caustic potash

On adding caustic potesh and potassium iodide to a strong solution of tetrethylammonium hydroxide, a white crystalline mass of tetrethylammonium iodide is produced

The salts of tetrethylammonium are mostly crystallisable and readily soluble

Twis-shiplanmonnum Chloride, (G_1H_2) , KG_1 , obtained by neutralising the hydro-and with hydro-dioric acid, is erystalline and highly daliquescent. It forms double solisivith auric, moveme, and platine chlorides. Tet-chiplanmonnum chloropdistrates, $(B_2K)_1^{\rm PLC}G_0^{\rm p}$ is thrown down mmediately as an enuge-yellow precapitate, or astupped platine chloride as solition of tetretilylammonium chloride. It is slightly solithle in water, and less solithle in alcohol and other

O collec and Schayver (Jone Chem See, 1vn. 767) have recently shown that when a nuced quantanay a monnoum collected or hydroxide (made from trunchlylamme or trachylamme) is heated, a mixed 1 set in its ry am in a diverge a chicked in greater or less amount. With truphanylmathylammenum the only preduct is dimethylphenylamne, whole with the allyls and garpept-transcriptammenum composeds, the chart tautry amme formed by the setting of the setting the setting transcriptament of the set of the setting true of the setting transcriptament of the setting tr

HYDRAZINES.

The name hydrazine was first applied by E Fischer to a hypothetical base, having the constitution of diamidogen, Hay NNL₂ Since then the base itself has been obtained in the form of a hydrate, and possibly also in the free state.

Hydrazine. Diamidogen. Diamide. NoH4 or H2N NH2. Hydrazine is obtained by the decomposition of tirazo-acetic acid by heating it with water or nimeral acids, when the following reaching occurs —

$$C_8H_3N_0(COOH)_3 + 6H_2O = 3N_2H_4 + 3C_9H_2O_4$$
.
Trace acets acid Water first again to the acid

The exalic and is more or less split up, according to the temperature and the strength of the and employed, into exploine and forme ands, so that when only water is used the hydranic separates as a formate, but if a mineral acid be present it forms the corresponding said.

Hydrama has an extraordinary affluity for water, acadity forming a hydrata, NJL₃H₁O, which it does also when set free from its safe by caustic alkales or line.¹ This hydrate is a highed funning on the an and being unileveral at 119 °C, and can be easily separated from water by distillation, though some of it passes over with the steam. When lacked with braight oxide in a scaled tube to 170°, some amply how it hydratine appears to be formed and escence so what the queen on conjunct the hybrid of the control of the configuration of the control of the

The solution of hydrazine tune reddened litinus-paper a deep blue, and gives white fumes with and yaponis. In a concentrated state it has a very peculiar odour, only slightly resembling that of

² Hydradne hydrade is best prepaised (Curtius and Schultz) by distilling armitter of eleven parts of hydradne shiplate with four of contributions armitter of eleven parts of hydradne shiplate with four of contain poised and one of water me anive retort provided with a suiver condiner. When the back drop has passed over, the distillates the Returnited. After four fractonzatous file lest portions bed constantly at 119° Out it us and 1 sy (fose Pract Chine, [2], xxxxx 27) peaper hydradne hydrating the hydrochlorade of the base owth carette time on a silver retori, and preveng the vygours though a hasted aliver their containing caused line.

ammona It powerfully affects the nose and throat, has an alkaline taste, and leaves a burning sensation on the tongua. When boiling, the solution attacks glass, and quickly destroys corks and india-rubber Hydrazine, like hydroxylamine, is a strong poison of universal character.

Hydiazune reduces Wehling's solution and ammonu-nutrate of silver in the cold. With cupric sulphate it yields a red precipitate (1 cuprous oxide), with mercuric chloride a white precipitate, and precipitates alimning from a solution of alium. With aromatic aldehydes and ketonos it yields sparingly soluble ciystalline compounds

SAUTS OF HYDRAZINE

Hydrazane combanes with one or two molecules of monobase acide to form very stable saits, which are usually crystalline and asmorphous with the corresponding ammonium salts. The salts Hz,2HR crystallies in the regular system and are resaltly soluble in water, but nearly insoluble in atchief. The mono-acid salts, HaHR, are easily soluble in water and waim alcohol, from which they crystallies well. The salts of both classes are moslible in ether, benzene, &c. In acid solution, the salts of hydrazane possess remarkably strong reducing properties, and are powefully toxic towards the lower organisms. Peytone solutions containing 0.1 per cent of hydrazine sulphate are unable to support bacerial life.

High anne Dhhytrochloride, N. H., 2RCI, cystallises from howers in large glassy octshedra that are freely soluble in water, but less so in alcohol. On treatment with platime chloride vidose not yield a chloroplatinate, but is decomposed with evolution of much nitrogen. It mells at 198° C, with evolution of hydrochloride, and, to a clear glass consisting of the momolydrochloride, N₂H₂HCI, and this on further heating to 240° C. is decomposed into ammonium chloride, nitrogen, and hydrocent.

High-name Sulphate, N.H. H.SO, according to T Curtus, is best obtained from ethyl diazo-acetate, which on treatment with hot concentrated caustic potash yields the potasium salt of an acid which separates in golden yellow lablets on addition of a mineral acid. On dispeting the yellow algueous solution of these with very dilute sulphure acid, the colour disappears without crolution of gas, and on cooling crystals of the sparingly soluble hydrazme sulphate are obtained. From the sulphate, other salts of hydrazme may be prepared by double decomposition with barum salts

Salts of hydrazine in solution are decomposed by sodium nitrite, with evolution of gas attended by much frothing. The reaction is analogous to the decomposition of ammonia salts by a nitrite, with the difference that whereas in the latter case (a) nitrogen is formed, in the case of hydrazine (b) azoimide, HN_{g_1} is found among the products of the leaction —

- (a) $NH_{sp}HCl + NaNO_{2} = NaCl + 2H_{2}O + N_{3}$.
- (b) N_2H_4 HCl+ $NaNO_2 \approx NaCl+2H_2O+IIN_3$.

AZOIMIDE IMIDAZOIC ACID HN3=IIN<

The above reaction is not a suitable one for the preparation of this remnskable body, which, according to its discoverage for t of the results of t of the results of t of the results of t of t of the results of t of t

NHDz CH₂ CO N(NO) NH₂ + 2NaHO = NHDz CH₂ COONa + 2H₂O + NaN₂.

On distilling the compound NaN, with dulute sulplume scal, mudazone acid volatiless with the steam, which when passed into a neutral solution of minute of aliver gives a precentate of the airer sult. Thus is weshed and decomposed by dilute sulplume acid, the solution being used instead of sulver initiate to absorb the vapours of mindazone acid. By repeating this process, a solution containing 27 rice ren't of the new arid is obtainable.

In the anhydrous state, unidazoic acid is a colourless gas of a peculiarly nauseous odom, and condensable on cooling to an extremely explosive liquid. It is very soluble in water, and on distillation of the liquid a concentrated acid passes over. the distillate gradually becoming weaker until an acid of constant composition and boiling-point distils. The solution reddens litmus, and gives white fumes with ammonia, of the salt NH. HN. or N.H., which sublimes completely at 100° C, but does not crystallise in the cubic system like ammonium chloride Iron, zme, copper, alummum and magnesium dissolve readily in dilute unidazoic acid (7 per cent) with evolution of hydrogen, and gold is dissolved with formation of a red salt. The silver (AgNo) and mercurous salts of unidazone acid are insoluble, the former closely resembling silver chloride, but not blackening in the light. Both the salver and the mercurous salts are extraordinantly explosive, 0 001 gramme of the former indenting an iron plate on which it is heated to 250° Barium imidu.oate, BaN separates from concentrated solutions in short shining anhydrous crystals, which explode with a green flash when heated, or exposed to a strong green light. The solution of currie imidacoate deposits cuprous oxide on boiling. The free acid is bheated from any of the mindeazoates on treatment with dilute sulphunic acid. With concentrated sulphunic acid, the azominde is itself decomposed. Ethes of immedazoa acid have been prepared, phenyl innidazoate, PhN₂ being identical with the diazobeinzolimide previously described by Gricsal

SUBSTITUTED HYDRAZINES.

Hydranna is the parent of a large and impostent class of bases generally called hydrasines, one member of which, phenylhydiazine, (C₀H₂)HNNH₂, has proved, in the hands of E Fischer and others, a reagent of the highest importance, memicious recent syntheses in the sugar group having been effected through its aid. By replacing a second atom of hydrogen by (e) phenyl, secondary hydrasines may be obtained either symmetrical like hydrasines may be obtained either symmetrical like hydrasines may he obtained either symmetrical like hydrasines may he obtained either symmetrical like hydrasines may he obtained either symmetrical like hydrasines (C₀H₂)HNH(C₀H₂), or unsymmetrical like hydrasines (heavy hydrasines) in their power of reacting with he haloud salts of the alkyl adoesd (e.g., sthyl-iodide) to form hydrasinism compounds.

 $R_2N NH_3 + AkI = IAkR_2N NH_2$

The hydraunes contaming fatty sikyl radicals are liquids boiling without decomposition; these of the assumatio series are resultly fusible solids or oily liquids, and are partially decomposed on distillation. Hydraunes itself and some of the fatty donvatives are di-acid bases, but the hydraunes of the benzene series have all monobases functions

The hydrazines closely resemble the amines, but are dis-

1 From the assertatived characters of mudanous and, and its analogy to hydrocymno call, Mondelged has formulated some very interesting prognestications. Just as ammonium formate, when heated, yadds formamidand the intrifie HON, so ammonium intend-decomposes on beating with production of (on intermediate hypedhetical intramide and) the intrile N₂O, introse oxide.

Similarly, azomude may be regarded as the nature of diaminonium orthonitrate, thus ---

Feimate, . . HCO O.NH₄ — 2H₂O = HC N , hydrocyame acid Meta-intrate, . O.NO O.NH₄ — 2H₂O = NO N , introus oxide Ortho-intrate, HO.NO (O.NH₄)₂ - 4H₂O = HN N₂ , inidazore acid

It seems not improbable that the ammonium sait of immission and, NH, HN, may prove convertiblement is symmetrical immensels, NH, MH, NH, NH, thin infinited friam min on rum or the on it rate, NO(ONH,)(ONH,)(ONH,), just as ammonium symmetre can be changed into use. The existence of explosive, coloured, double invidence is forested by Mendeleying.

inguished from the latter by their capacity of reducing Febling's copper solution, in many instances at the ordinary temperature. The product of the oxidation of the hydranine is the corresponding amine Thus, dictilled 1 in dictilled 1 in dictilled 2, dictilled 3, dictilled 3, dictilled 3, dictilled 4, d

The general and special characters of the hydrazines are sufficiently exemplified by two typical species, ethyl-hydrazine and phonyl-hydrazine

Ethyl-hydrazine, C₂H₂N₂=(C₂H₅)HN NH₄.

On treating diethyl-ures with introns and, a nitrosocompound is formed, which on reduction with rise-dust and seets and reconverted into a body called diethyl-some arbaride.

$$\begin{array}{cccc} \text{TNH}(C_0H_6) & \text{CO}\left\{\frac{NH(C_0H_6)}{NH(C_0H_6)} & \text{CO}\left\{\frac{NH(C_0H_6)}{N(NO)(C_0H_6)}\right\} & \text{CO}\left\{\frac{NII(C_0H_6)}{N(NH_c)(C_0H_6)}\right\} \\ \text{Diethyl-meta} & \text{Nili opo compound} & \text{Diethyl semicorbasics.} \end{array}$$

This last body decomposes, on heating with strong hydrochloric acid, into ethyl-hydrazine, ethylamine, and earbon droxide --

$$\begin{split} \mathrm{NH}(\mathrm{C}_2\mathrm{H}_5) & = \mathrm{CO} \ \mathrm{N}(\mathrm{NH}_2)\mathrm{C}_2\mathrm{H}_5 + \mathrm{H}_2\mathrm{O} = \mathrm{HN} \ (\mathrm{NH}_2)(\mathrm{C}_2\mathrm{H}_5) + \\ & = \mathrm{HN} \ \mathrm{H}(\mathrm{C}_2\mathrm{H}_5) + \mathrm{CO}_2 \end{split}$$

The ethylhydrazme hydrochloride is less soluble than the corresponding salt of ethylamme, and may be separated from it by crystallisation

Ethylhydranae is a colourless, mobile liquid of ethered and family ammoniacal odon. It boils at 100°, and distits underomposed. It is very hygroscopic, forming white funes with mostan, dissolves in water and alcohol with evolution of heat, and corrodes cotk and contchoine.

Ethylhydrazine gives Hofmann's isometrile reaction for primary amines with chloroform and alcoholic potash (1920 7). Bromino decomposes it with evolution of nitrogen, and it is also decomposed by introgen broade.

Ethyllydramme is a very powerful devotdsang agent. It reduces feeling's copper solution at the unimary temperature, reduces asgentic oxide, and converts evade of unerony into mercurities of thirds, $\mathrm{Hg}(\mathrm{C_2H_2})_2$. It yields a black precipitate with Newder's solution.

Ethyllydrazme reacts with aldehydes, with evolution of heat, to form ethyl-hydrazides, RCH-N2H(CoHz)

Potassium anhydrosulphite, K₂S₂O₇, reacts on ethylhydrazine to foim potassium ethyl-hydrazine sulphite,

(C.H.)HN.NH(SO.K),

which, on treatment with mercuric exide, gives potassium diazo-ethane-sulphonate, C_3H_6 N N (SO₆K), a substance which explodes violently when warmed, and otherwise resembles the diazo-benzene-sulphonates (Part I page 187)

DIRECTIFIED ALL MINISTERS AND ALL MAN THE ALL MAN ALL

Phenyl-hydrazine. C,H,N,=(C,H,)HN NH.

Phenylhydrianne is prepared by the action of reducing agents on dia zo be nzene compounds, C₆H₆N NX (Pait I page 176) Thus diazobenzene chloride may be reduced by the calculated amount of stannous chloride and hydrochloric acid, or the potassionallylate with zinc-dust and acotic send, the product being subsequently decomposed by boling with hydrochloric acid.

 $C_6H_5HNNHSO_8K+HCl+H_9O=KHSO_4+C_6H_5HNNH_9,HCl^1$

Phenylhydrazine is a yellow oil of a faint aiomatic odou. It solidifies at low temperatures to a crystalline mass, melts at 23°, and boils, with slight change and evolution of ammonia, at 241°-242°

¹ Picuylih drame a best obsumed, as described by Y Me y rs, by dasslving 1000 parts of anims in 2000 parts of anims in 2000 parts of anims in 2000 parts of summin series of strong hydrochloure and, cooling the solution by means of nee, and thes slowly adding an nee cold solution of 10 reputs of solution mutter and 00 c of water. To the cold solution of 10 parts of series can be near the cold solution of 10 parts of series can be near the cold solution of 10 parts of series can be needed to 10 parts of 10 parts o

It volatilises in a current of steam, but not very readily Phonylbydrazine dissolves sparingly in cold water, more readily in hot, and very readily in alcohol, other, chloroform, and benzenc

Phenythydrazme is readily oxidisable, and becomes red and ultimately dark brown on exposure to air, from absorption of oxygen.

Phenylhydrazme has well-marked antiseptic properties, and a 0-1 per cent solution of the hydrochloride has been accommended as a substitute for one of mercuric chlorido of equal strength (Therm Josa. [31, xix 608).

Under certain undetermined conditions, contact of phenylhydrazine with the skin produces troublesome serve

Phonyllydranae has well-matked brase properties, and forms well-crystallased salts. The hydrocollorate, prepared as already described, crystallases from but water msmall, thun, tustous pattery, and as almost completely precipitated from tax anyeons solution by concentrated hydrocollora card, a reaction by which plenyllydranae may be roadily spanated from annium and sevend other bases.

Solutions of the hydrochlorudo and other salts of plenyllydrazine act as powerful reducing agents They reduce the salts of silver, mercury, gold, and platinum in the cold Freshly-necipitated mercuric oxido is reduced, a salt of diazohenzene being reproduced Echling's solution is reduced in the cold, with evolution of integen and precipitation of curpous oxide, annino and benzene being

hydrochloude be treated with a cold solution of potassum nitude, a nitroso-con pound, C₄f₄f₄(NO)N Ni₂, separates in yellow flocks, which, on treatment with phenol and strong sulphune acid, yield a hown solution, changing to given and blue. This socion, observed by Liebermania, is common to all miscos-forwalves.

Phonylhydrazino combines directly with carbon disable, carbon disablehide, and cyanogen The sulphonic acid (paia) is employed for the preparation of tartiazia (Part II page 288) and other dves

Prinkythytomators. The acctyl-derivative of phenylhythuraue, C₂II₂ Hin.NII(C₂II₃O), which may be regarded as a c e t + ph on y t + by d r a z i d, has powerful antipyrebe properties, and has been introduced into Genman pharmacy under the name of "hythrectin". The same substance is said to be the active impreduct of the preparation known as "py i od in e" (Pharm. Journ. [3], xix 425, 508, 1049). Both substances seem to be uncertain in their action and dangerous in use; in fact, hydraceth is reported by Ke in vers to be a direct blood-poson, the antithermic properties of which are ionly due to destruction of the red corporates.

"Orthine" is the name given by R Kobert to a body having the constitution of an orthohydrazine-parahydroxybenzoic acid .--

The free base is very unstable; but the hydrochloride is stable, reduces the porealts of the heavy metals, and possessee a marked authorite action

Phonylhydrazine in aqueous solution reacts very readily with the hydroxy-acids of the suga $^{\prime}$ and galactomic acids, $C_8H_6(OH)_8COOH$, $C_0H_{18}O_7\rangle$

with elimination of wites, to form crystalline phenylhydiasales, RCO HN NH(C₁H₂). They are prepared by treating a 10 per cent: solution of the and or its lactione with a modernic access of phenylhydranne and an equal quantity of 50 per cent sectic and, and heating the mixture to 100° for 80 to 120 minutes. The hydrazide sometimes crystallines from the hot solution, but more usually by social about 50 minutes. The hydrazide sometimes crystallines from the hot solution, but more usually by social about 50 minutes. The hydrazide by social about 50 minutes with the solution of the high solution of the hydrazide by the solution of the hydrazide by the hydrazide by the hydrazide by the hydrazide by crystallinetic from hot water. The products are beautifully crystallines, those derived from monobasic acids being but little solution in cold, and only with difficulty solution in the water, while those from polybasic 1 and mixed) are still less results.

isomeric acide usually present a close issemblance in their physical properties, but the acide from which they are derived can be regemented (in a pure state) by boiling the hydratide for half an hour with thirty volumes of 10 per cent baryta water, which treatment hydrolyses them completely. From the product, the phenyi-hydratine is extracted by agitation with ether, and the aqueous liquid, with any precipitate which may have been formed, is boiled and treated with sulphuric seed in quantity sufficient to precipitate which burnum as BaSO. The filtered liquid yields the free and or lactone on evaporation (Fischer and Passmore, Ber, xxii 2728, Jose, Chem Soc. 1 vin 152)

The hydrazides are coloniless and readily hydrolysed by alkalies and bayta They can be readily distinguished from the hydrazones by the reddish violet coloration they give when dissolved in strong sulphinic soid and treated with a drop of ferric chloride solution Hydramores. Phenylhydrame behaves in a lightly interesting manner with bothes having the constitution of aldehydes and ketones, with which it seeds with elimination of water to form compounds called hydramore of the Most of the bothes of this class are sold and crystalline, and therefore well suited for the recognition of the aldehydes or ketones producing them. The reaction appears to be general for bothese containing the carbon vil group, CO. The reaction is sometimes complicated by the presence of other reactive groups. Thus compounds containing the arks to neal to the other containing the arks to neal to the containing the group—CH(OH) CO—, next in the cold with only one molecule of phenylhydrame to form colouriess compounds containing the group—CH(OH) (N MCA),—

Observation where the compound thus formed is heated with excess of phenyllybrianns, the alcohol group undergone thelydic-genisation, is eaking at the same time with a second molecule of phenyllydramae and gring rise to a relief compound containing the complex group—C(N.NIII.G.I₂) C(N.NII.G.I₃).— Compounds of this kind, in which two bydramae-residues are attached to two contgeones can bon-atoms, are called to a s.c. on a, and maybe obtained directly by the action of phenyllydramae on the di-ketones. They are of interest in commentor with the carboliylates, which may frequently be recognised by means of their characteristic examines of the control of the

Pyrazotives An unsaturated hydrocarbon group (r.q., allyl, C_SH₆), if contiguous to the carbonyl group, may also react with phenylhydrazine —

Pyrazolones.

The pyrazolones are derivatives of a body of the formula $C_1H_1 N_2O_2$, the synthesis of which has been effected by Balbiano ($\hat{B}er$, xxiii 1103). The relationship of pyrazoline to pyinzol, pyrazoline, and pyrazoline is shown by the following formula::—

$$\begin{array}{l} \text{Ppracel} \\ \text{N CH} \\ \left\{ \begin{array}{l} \text{N CH} \\ \text{CH} \end{array} \right\} \\ \text{HN} \\ \left\{ \begin{array}{l} \text{N CH} \\ \text{CH}_2 \\ \text{CH}_2 \end{array} \right\} \\ \end{array} \\ \begin{array}{l} \text{IIN} \\ \left\{ \begin{array}{l} \text{N CII} \\ \text{CO CHI}_2 \end{array} \right\} \\ \text{IIN} \\ \left\{ \begin{array}{l} \text{NII}_1 \\ \text{CII}_2 \\ \text{CII}_2 \end{array} \right\} \\ \end{array}$$

This body must not be confounded with placing, which was formuly called pyrazing, and probably has the constitution :-

(See A. T. Mason, Jour Chem Soc. ly 97.)

Phenyl-Pyrazolone, C_6H_5 $C_5H_5N_2O$, is obtained by heating phenylhydrazine and iodopropionic and together to 100°, and treating the product, in chloroform solution, with mercuric oxide.

PHENYL-METHYLPYRAZOLONE, CucHioNaO,

$$\begin{array}{ccc} & N \ C_6H_5 \\ N & CO & , \ \mathrm{or} \ C_6H_5 \ N \left\{ \begin{array}{c} N \ C(CH_2) \\ CO \ CH_2 \end{array} \right\} \end{array}$$

When phenylhydramae is added to thy lie actio-actite, CH_{S} CO CH_{S} CO CC_{S} Lb, the two bodies react in the cold, with elimination of water, to form CH_{S} C(N $\mathrm{NHPh})\mathrm{CH}_{S}$ CO, CC_{S} Lb). On heating, the hydrixone thus formed splits up into alcohol and phenyl-methylpyiazolone, a body which was originally regarded by its discoverer, Knorr, as a methyl-oxyquinizine

To prepute pheapyl-methylpyrazolone, 100 pairs of phanyl-hydrame are added to 125 of ethyl neot-cactes, the water which forme is separated, and the only product is heated for two hours on a water-bath, until a portion is found to colvidity on cooling, or on the addition of ether. The warm mass is pound into and shired with ether, which removes colouring matter, and the wint crystalline product washed with ether, and dred at 100°. The yield is quantitative and the product pure. It is almost insoluble in cold water, ether, and petroleum spirit, more readily in hot water, ander, and castly in alcohol. It crystallises from hot water or alcohol in hard builting prisms. The hydrochurde, O_wH₀N, O_wOHCi + H_OO, males

ANTITITIBMN When an squeous solution of levulina cost (sosto-promosous and), OR OCH, CH₂ GO, OCH, a sadded to an equivalent amount of baselylivetzame, dissolved in dilute sceine said, a yellow productiate as produced of the hyd at so. so. a CH₂GON MPRIO (Mr. GH, COCH). When recrystated from alcolod, thus body forms large colouriess, odcuriess or yatalo of a slight butter test, mairing at 8°-89°, and meanly would hid on which the state, the solid is a 18°-89°, and could be used to the solid of the sol

3 Whan a number of placylmosthyl-gyrandome and phorphydrames is meated to boung displacely, methyl plys as lone, 62H₃N₂O_c, as formed. Healed with methyl obcoled or methyl nobies it yields disnet, formed. Healed with methyl obcoled or methyl nobies it yields disnet, prince, 62H₃N₂O_c, mething at 98H₃O_c and datengualed from antipyrum byte sparing solubility in vates and the melting-point of its peasts (161°) When the body C₃H₃N₃O_c, in restend in allatime colution with excess of solution untrite, and the mething-point of the subdime soid, promod-form billion and the mething before the contribution of the contributio

at 96°, and the chloroplatinate, $(C_{10}H_{10}N_2O)_{+}H_2PtCl_{+}+4H_{*}O$, in presses melting at 110° Phenyl-melhylpyrazolone yields crystaline preceptates with salts of many of the heavy metals. With silver intrate an aqueous solution gives crystals of $C_{10}H_{5}AgN_{*}O+C_{6}H_{10}N_{*}O$. The ultramarine cobalt compound and the orange-yellow uranium salt are especially characteristic

PHENYL-DIMETHYLPYRAZOLONE ANTIPYRINE PHENAZONE

$$\begin{split} &C_{11}H_{12}N_2O = C_3N_2H(C_6H_6)(CH_3)_2O & [Ph.\,M_6.\,M_6 = 1:2:3:3];^1\\ ∨,\,CO\left\{ \begin{array}{c} CH\cdot CMe\\ NPh.\,NM_6 \end{array} \right\}; \;or,\,C_6H_5,N.\left\{ \begin{array}{c} N(CH_7).C(CH_7)\\ CO - CH \end{array} \right\} \end{split}$$

When phenyl-methylpyrazolone is heafed with methyl solide, or further substitution fakes place, with formation of phenyl-dimethylpyrazolone, a cabstone known geneally as "antipyrine," isocommonly se "an i gesin," and called in the additions to the British Pharmacoppias (1890), phenorone It is official in the Ge man Pharmacoppias of 1890 under the name of Autrypy into

Antipyrme is prepared by heating equal parts of phonyl-mothylpyrazolon, suchlyl oddie, and methyl alcohol to 100° in a closed vessel. The dark product is decoloused by bothing with sulphurous acid, the alcohol distilled off, and the residue shadem with strong soda, when the base separates as a heavy oil. This is separating and treated with either, in which it is spannigly soluble. On separating the other and evaporating off the solvent, the antipyrme is obtained as a mass of crystals which are purified by recrystallication from tolucers.

Antipyrme forms small, lushous, rhombes needles or plates, which are colouless, but have a converbal tolter taste. When parfectly anhydrous it melts at 110° to 113° (BP, 110° , GP, 113°), but on exposure to are takes up a smill proportion (0.6 pc ont) of waker, and in that state melts at 103° – 103° – 103° . Of . The hygrescopic water may be driven off by exposing the substance to a temperature of 100° , when the original melting-point is restored.

a temperature or 100°, when the original metang-point is restored.

Antipyrine is soluble in about its own weight of cold water, and
in less than half its weight of bothing water. It theselves in twice

acid has an indigo blue colour, and gives an absorption spectrum resembling that of indigo. It is not a substantive dye, is decomposed by strong all thes, decolorised by chlorine and intric acid, and converted into drephenyl-methylpyrazolone by reducing agents.

² Two asonats of anthyrme have been pregated, and others are critile existing. The known asonas differ from anthyrme by length as soluble in water, not yielding into so-detivatives, and by giving methyr audine either when distilled with anne dust or heated with hydrochloric aid to 200° under pressure.

tts weight of absolute alcohol, but in little more than its own musch of rectified spirt. Anthrpynne is soluble m an equal weight of cathyle alcohol, and m one and a half times its weight of chloro-form, but requires about fifty parts of either for solution, difficultly soluble in benzene, and nearly insoluble in petioleum synth.

On adding strong caustic soda to an aqueous solution of antinyrme, the base separates as a milky precipitate, which speeduly collects into only globules. On adding a little either, these immediately solidify to white crystals without appreciably dissolving, but they dissolve metantity on adding shloroform (J C W at e t o u s e)

An aqueous solution of antipyrine exhibits no alkaline reaction with litimus or phenol-phthaden, but destroys the red color of an acidulated solution of methyl-orange. Free antipyrine may be determined with accuracy by titration in aqueous or alcoholic solution with methyl-orange.

Antipyrne is a strong monovalent base Its sails, most of which are solutible, do not readily crystalles, with the exception of the preats (molting at 188°); the percognants (C₁₁H₁₂N₂O₃), H₁Cfy, which forms a crystalline preceptate; the chlorophatinate, (C₁₁H₁₃N₃O₃H₂PCI₃+2H₂O, which forms rellevishered pressure melting at about 200°, and the exheptate (page 37)

When antipyrno is heated with hydrochloue and under pressure to 200°, it suffers complete decomposition, yielding much a niline and a small quantity of m sthylamine, besides other products On distillation with zime-dust it yields benzone, aniline, a bease boiling at 86°-87°, and other products

Antipyrine is unchanged by treatment with reducing agents in the wet way, but with oxidinary agents it gives a sense of interesting reactions (G ay and Fortuné, Pharm Jour, [3], xvincibles in the when boiled with potassium chlorate and hydrochlorie setd, antipyrine gives a reddsh-yellow liquid, which on cooling deposits brightered only globules, site up by chloraform with greenish-yellow colour. A solution of bleaching powder produces no chinge in the cold, but on hesting a broke-wed precipitate, is formed, and the highest coloured yellow. Sodium hypochlorite is add to give the yellow colourion on hesting, without any precipitate being formed. Chlorine-water produces no change, and bronne-water a light yellow precipitate, assolving on hesting Potassium bichromate and permanganate are reduced by acid solutions of antipyrine.

When a solution of iodine in iodide of potassium is added to a solution of antipyrine, a precipitate is formed which disappears on agitation, leaving the solution colourless, but on further addition of the reagent a permanent brick-red precipitate is produced, perceptible in a dilution of 1 in 20,000. According to Manseau (Pharm. Jow , [3], xx 162), the point at which a permanent precipitate is formed is perfectly definite, and he suggests that the purity of a sample can be ascertained by titration with a standard solution of iodine Millard and Stark (Pharm. Jour, [3], xx 863) find that the point of permanent precipitation depends to a marked degree on the dilution of the antipyrine solution. Thus in a 1 per cent solution, 1 gramme of antipyrine gives a permanent precipitate after the addition of 39 c.c. of decinormal iodine, while with twice the volume of water 7.2 cc. are required. The authors state that more concordant results are obtainable by using starch as an indicator of the end of the re-They dissolve 0.5 gramme of the sample of antipyrine in 200 cc, of water, add plenty of starch solution, and then drop in decinormal iodine solution gradually until a distinct blue coloration is obtained, which does not disappear on vigorously shaking or stirring the mixture. E Munzer has described an indo-antipurine, C., H., IN.O, which forms colourless, tasteless needles, meltmg at 160°

An acad solution of mercure nitrate gives a white precipitate with a solution of antipyring 2 co of Milliur's region at ad 4 cc of a 1 per cent (noutral) solution of antipyrine give a white precipitate in a yellow picular, in a solution and with hydrochloric scal, a yellow precipitate in an orange-yellow liquid, in a solution and with hydrochloric scal, a yellow precipitate in an orange-yellow liquid, in solution ten times more dilute a yellow precipitate and green liquid result, and in an acid solution of part of antipyrine in 20,000, a white precipitate and yellow liquid 1 cc of a saturated solution of functions missio added to twice its measure of a 1 per cent solution of antipyrine gives a yellow precipitate floating on a bloot-yed liquid

If antipyino be heated with strong ritric and till reaction commences, and the liquid be then allowed to cool, a fine purple coloration is produced; on adding water a violet precipitate is thrown down, and the filtered liquid is purple-red

Nitroscaningrino Several of the foregoing mactions are probably due to the presence of intruss acid, which (if abled in the form of red funning mirro acid) gives with a 1 per cent solution of authyrino a heuriful green coloration, soil prorrightly when diluted to 1 in 20,000; when the liquid is heated it becomes purple red In strong solutions a copious formation of small, green, needle-shaped crystals occurs. These consist of isonitroscan, needle-shaped crystals occurs. These consist of isonitroscan artipyrino, Gulfin, RNOIN, Q. and are best obstanced by adding a solution of solution intrito to a solution

of antipyrne in accidulated water. The liquid at once becomes blush green in colour, and an abundant formation of crystals specify occurs. These may be washed with cold water, and drod at the ordinary temperature. Nitrose-antipyrne explodes when heated to about 200°, is nearly mobilithe in water and ditties ands, soluble in alkatics and in aceta card, inderstally soluble in alcohol, and sparingly in chloroform and either By treatment with zinc and acetic acid it is converted into an oily because.

The green coloration of antipyrme with introus and is delicated and, to a certain extent, characteristic, but is common to all pyrazolones A C S tank recommends that the test should be applied by dissolving potasium intrite in a test-tube in a literal water, adding excess of strong sulphure and, and then filling the table with the limult to be tested.

Antipyrine dissolves without colour in pure anhydrous ethyl nitrite, but a green colour is immediately developed on addition of water When antipyrine is added to spirit of nitious ether containing free acid, the mixture iapidly acquires a dark-green tint, and green needles of nitioso-antipyrine separate, reaction (which does not occur if any free acid be neutralised by potassium bicarbonate) delives practical importance from the fact that spirit of nitrous ether and antipyrine are not infrequently dispensed in conjunction. A mixture of the kind is alleged to have been fatal to the patient, but it is very doubtful if the nitroso-delivative of antipyrine was the cause of death, for direct exhibition of the compound to a small rabbit, both hypodermically and by the stomach, in doses commencing at 1 gram, and gradually increased to 4 grams, produced no perceptible toxic effect (Pharm Jour, [3], xviii 1085) Similar experiments have been made on dogs (Pharm. Jour., [3], xix. 807)

Anthynine gives a very delicate and characteristic reaction with ferric cilorite, which, in a 1 per cent solution, produces a blood-red coloration. The reaction is still very distinct in a solution of 1 in 2000, and perceptible at a dilution of 1 in 00,000. The red coloration is destroyed by excess of mineral acids. The reaction is at once given by urine containing antinyrine.

On mixing cold aqueous solutions of antipyrine and mercuric

² The haud filtered from the crystals gradually changes colour from green to hown, and after standing for some hours as found to smell of hydrocyanio acid, but the quantity of this body formed appears to be very mutuic (Wood and Maishall, Pharm Jow, 73), xix. 806)

chloride, a white precapitate is formed. On boiling the liquid this disappears, but on continued boiling a brown resulted substance is deposited, which, when separated, is found to be soluble in hot alcohol and in nitric and, and is colouned scarlet by strong subburne act.

Antapyrme seeds in the general namum of alkalouds. Thus, in and solutions it gives a yellowish-white promptate with Maye's reagent, and the same with Marind's test (polarso-cadmum nodide); a green precipitate changing to orange-red with potasso-ordide of bismuth; an abundant reddish-y-rlow precipitate with Nessler's reagent; a white with phosphomolyhdate of sodum; and an abundant white precipitate with anum.

According to the German Pharmacopeau, the solution of antipyrine in two parts of water should be neutral, free from acrid taste, and not changed by sulphurested bystrogen water A 2 per cent solution should give a white precipitate with tainfin, and on addition of two drops of funing intree and to 2 c. o. of the solution, a green coloration should occup changed to red on boiling and adding another drop of intree and. 2 c. of a 0.9 per cent solution gives a deep red colora with a drop of ferric chloride solution, changed to bright yellow on adding 10 drops of sulphure and Sunta twist are given in the additions (1890) to the British Pharmacopeaia, in which antipyrine receives the designation "fisherson".

Anthyrme has now an established position and wide apphication in motioner. Although originally intioduced as a febringe, it is taking a still higher place as an anodyne Given in 10 to 20 gram dosses in cases of bittons and nervous houshche, it often effects a remarkably rapid and perfect one. It has been usefully myected hypodermosily in 8-gram dosse as a substitute for morphia, and for the relief of pain in acute and chronic gout, neuralgia, sensitian, &c. The obsertanceous injection of antipyrine is said not to be followed by drowniness, conting, or excitament. It is stated to be almost a specific in pureprisal fever. It has been found valuable as a thomostatic, and has proved successful in some cases of sea-suckness, but by no means invariably. Antipyrine causes an iduost inniculate re-

³ The reactions described in the text sufficiently inducte the pharamer-interpreparations with which attriprise is incentiable. Their it should not be dispensed in a maximir with intre scal, mitries, chloral hydrict, solid solium, assignistic, archieb scal, famin, before, meneure chlindric, saits of home permanguastes, or timetures or influence of cateclin, embions, roces, galls, rimbarb, & or with 111s at and 81 start, */Ram* Jan* (3], 3x × 800).

² Autipyrine has been adulturated with acetanihile (see page 72).

duction in the temperature of the body (apparently) from its influence on the brain-centres royulating the temperature), the effect continuing from four to six hours. It induces sweating and feeble pulse, and in excessive doses, or even small doses in certain cases, an eruption resembling nettle-isal, occasionally with vomiting and collapse. Atropine has been found to act promptly as an anticlote.

Antipyrne may be detected in the urne for eighteen to twenty-four hours after it is taken by the atomach, but can detected only for a few hours in the different organs. It has been detected, after purefaction for a fortingit, in animals life within two hours after its administration, either by the stomach or lyrodermucally.

Antipyrine is readily extracted from animal matters, by rendering the hquid ammoniacal and agitating it with chloroform or amylic alcohol.

Antipyrine Salicylate, C11H10N0O, C7H0O0. If salicylic acid be gradually added to a dilute boiling solution of antinyrine, antipyrine salicylate separates as a yellowish oil The compound can be more conveniently prepared by heating equivalent proportions of antinyrine and salicylic acid with a little water to 90°, or by shaking together an aqueous solution of antipyrine with an ethereal solution of salicylic acid, when the salt separates in fine crystals Antipyrine salicylate melts at 89°-90° C, and decomposes at a somewhat higher temperature, dissolves in 250 parts of cold water more freely in hot, and readily in alcohol, ether, chloroform, and carbon disulphide. The aqueous solution is faintly acid in reaction, and has a sweet taste and bitter after-taste. It gives a violet coloration with ferric chloride, and green with nitrous acid. Salicylate of antinyume has been employed with favourable results in medicine under the name of "salipyrin" A mixture of antipyrine and saliculate of sodium gradually changes to an oily liquid on exnosure to air. The change, which does not occur in a closed bottle, appears to be simply due to absorption of moisture by the salicylate and the solution of the antipyrine in the water thus absorbed.

Antipyrune becomes pasty when mixed with betamphthol, and appears to form a compound with phenol. Under the name of "resopyrin," Portes has described a compound obtained by mixing solutions of molecular proportions of resoreinol and antipyrine. It crystallises in oblique rhombic prams, insoluble in water but solution alcohol.

1 The exhibition of antipyine is unsafe when the heart is week. A case where sowers symptoms were produced by a dose of 1 gramme has been recorded by Schwabe (Pharm. Jou., [3], xx. 1059).

Chloral-Autopoine, C., H.11(C.H.C.R.) NI,O. When diute solutions of chloral hydrate and antisyrne are maxed no perceptible reaction occurs, but on concentrating the hquid, or on mixing stong solutions of the two substances, as operation of only globulor takes places, and these immediately or gradually shange to a mass of crystals of chloral-antisyrne. The same substance may be obtained by heating molecular proportions of chloral hydrate (185 parts), and antisyrme (188 parts) to 110°-115° C. The reaction consists on elimination of waker and substitution of the group CCl. CH(OH) for one of the hydrogen atoms of the antisyrme, ¹ but whether the replaced atom so one of those of the methyl groups, or the hydrogen atom of the methyl groups, or the hydrogen atom of the first place of the methyl groups, or the hydrogen atom of the CH group, is not definitely decided (compare Phan m Jour. 181 xx news 889 with reage 889).

Chloral-antipyline, also ealled h v p n a l. civstallises from alcohol in hard scales and from water in transparent rhombs. It melts at 67°-68°, is almost odomless, and has a siline taste with an aftertaste suggestive of chloral It is only slightly soluble in cold alcohol, ether, and chloroform, but somewhat more soluble in boiling alcohol, and is dissolved by about cight parts of warm water The solution reduces Fehling's solution on warming, gives the blood-red reaction of antipyrine with ferrie chloride, and yields chloroform when heated with dilute caustic alkali. When chloralantipyrine is kept in a melted state for some time, it deposits ervstals of a dehydration compound, which is insoluble in water. melts at 186°-187°, and gives no colour-reaction with ferric chloude According to Reuter (Pharm. Jour. [3]. xx 602) chloral-antipyrine is physiologically mert, but Bardet found doses of 1 gramme to induce sleep as readily as chloral hydrate, while in cases of insomnia caused by pain it seemed to have the same anodyne effect as antipyrine Sehmidt finds the monochloralderivative to have more decided soporific effect and a less deleterious influence on the circulation than antipyrine.

Bishloral-Antaparnes us obtained by heating antipyine with excess of a strong solution of chloral hydrate, when an only species is formed, which solutifies to pusmate crystals melting at 67°–68°, soluble with some dissociation in ten parts of sold water, and giving the reactions of chloral-antipyrine

Butyl-chloral behaves similarly with antipyrine,

BASES FROM TAR.

The numerous constituents of tais may be roughly divided into-(a) Indifferent Bodies -as Hydrocarbons,

- (b) Acid Bodies .- as Phenoloids and Acetic Acid; and (c) Bases -as Ammonia, Aniline, Pyridine, &c.

The principal members of the first two groups have already been considered at length Ammonia is beyond the scope of present work, and the remaining bases which require consideration all belong to the aromatic group They may be arranged in several groups, each one of which is represented by a typical member

Thus ---1 Aniline, or Amido-benzene, CaΠ, NHo, or CNH. OR 2 Naphthylamine, or Amido-naphthalene, C10H7 NH2, or . . . 3 Pyridine, C5H5N, or 4. Quinoline, Coll, N, or OFF 5. Acridine, C, H,N, or

From these formula it appears that the substitution of nitrogen is outside the ring in the case of aniline and raphthylauline On the other hand, pyridine, quinoline, and aeritine are derived from henzene, naphthalene, and authracene respectively, by the substitution of N for one of the CH groups of the closed chain

Naphthylamine does not appear actually to exist in coal-tar, and, aniline occurs in tar in very limited quantity; these bases are obtained synthetically from constituents of coal-tar.

Besides the foregoing typical bases and thur allies and derivatives, certain volatile bases (eg, p) inpendine, conine, nicotine), ordinarily prepared from plants, and therefore classed with other vegetable alkaloids, have a connection with pyndine or quinoline which is now fully demonstrated.

ANILINE AND ITS ALLIES.

Annine is the type of a large number of organic compounds of synthetical origin

Anilme has the constitution of a mono-a mid obenzene or mono-phenylamine, and may be regarded as originating in the replacement of one of the hydrogen atoms of the benzenering by the group a mid og on, NH₂, or one of the hydrogen atoms of amonan by the radical phenyl, C_R||_L Thus.—

$$C_6H_5NH_2$$
, or $(C_6\Pi_5)$
 H
 H

Aniline exists in minute quantity in coal-tar, but is ordinarily produced by nitrofying benzene, C_0H_0 , and reducing the resultant nitrobenzene, $C_0H_0NO_2$, by nascent hydrogen.

If the treatment with nitre and be carried further, dinitrobenzene, $C_0H_4(NO_2)_2$, is produced, and this by reduction is converted into meta-phenylene-diamine or meta-diamido-benzene, $C_0H_4(NH_2)_2$.

If the reduction of mixebenieme be effected by alkaline magnetic, two molecules conclases, and a zobenson, $C_{\rm cL}_{\rm R}N$. $N_{\rm c}G_{\rm L}^{\rm H}$, is produced. On further treatment of this (especially mislonhoit solution) its converted into by drazoben zene $C_{\rm cL}^{\rm H}N$ is $N_{\rm c}N_{\rm c}^{\rm H}N$, which by intramolecular change is transformed into benadine or drapara-amido-diphenyl, $N_{\rm c}N_{\rm c}$ ($C_{\rm c}N_{\rm c}^{\rm H}N_{\rm c}^{\rm H}$

¹ Hydrazobenzene has no basic properties.

Analu e (Amidobenzene).	Andone	Analyse (Phenylamine)
NH.C.H.H	C ₆ H ₅ NH H	C ₆ H ₅ NH.H
Phenylone diamene	Phenylhydrazuse,	Diphenglamino
$NH_2C_0H_4NH_2$	C ₆ H ₅ NH NH ₂	C ₆ H ₅ NH C ₆ H ₅
Benndine	H ydrazobenzene 1	Hydracobenzene 1
NH_2 , C_6H_4 , C_8H_4 , NH_2	C_6H_5 NH,NH C_6H_5	C ₆ H ₅ NH.NH C ₆ H ₅
Aniline forms two class	sses of homologues	The true homo-
logues (Class A) coex	cist with aniline in co	al-tar, and are derived.
from aniline by the sub	estitution of one or m	ore methyl groups for
a corresponding number	er of the hydrogen a	toms of the benzene
analone There are and	demands absorbed by	mateurfrance the come

nucleus. They are ordinarily obtained by nitrofying the correaponding hydrocarbons prepared from coal-tar naphtha, and reducing the resultant nitro-derivetives. Thus -

Hydrocarbon Benzene—	Nitro-dermatice Nitrobenzene-	Amido dermotive Amline—
C_nH_nH	$C_nH_nNO_n$	C ₆ H ₆ NH
Toluenc-	Nitrotoluene-	Toluidine—
$C_nH_4(CH_n).H$	CeH4(CH2) NO2	C ₆ H ₄ (CH ₂) NH
Xylene-	Nitroxylene-	Xylıdıne—
CaHa(CHa)a H	$C_6H_3(CH_8)_2.NO_2$	Colls(CHs)2 NH
Cumene-	Nitrocumene—	Cumidine-
CaHa(CHa)a H	C.H.(CH.), NO.	CaHo(CIIa), NH
0 20 375	0 20 00 2	0 21 00

Isomeric modifications are known of all the members of the series except those in the first line (page 51 et seq)

The pseudo-homologues of aniline (Class B) are derived from anning by the replacement of one or both of the hydrogen atoms of the annido-group by methyl or other alkyl radical. Similar substitutions can be effected in the amido-groups of toluidine, xylidine, &c.

These alkylated anilmes (Class B) are obtained by the action of methyl chloride or other alkyl salt on annine, or of the corresponding alcohol on the hydrochloride or other salt of aniline (see page 73). Paratoluidine has also been obtained in a very interesting manner by heating the hydrochloride of methylaniline 2 to 350° C in a sealed tube, when change of position of the atoms within the molecule takes place thus -

$$\begin{pmatrix} C_8H_5 \\ CH_3 \\ H \end{pmatrix}N \approx \begin{pmatrix} C_6H_4(CH_3) \\ H \\ H \end{pmatrix}N$$

Mathel as the contribution

¹ Hydiazobenzene has no basic properties

² If the hydrodide of methyl amiline be similarly treated, ortho or metatoluidine is obtained.

By the same process methyl-toludine may be convexted into xylidine, and this by consecutive steps into a pseudo-cumidine, isodiuridine, and amido-pentamethylbenzene (page 60). By treating aniline hydrochloride with aniline, $d_1 p_1 = p_1 p_2 = p_2 p_3 = p_4 p_4 = p_4 p_5 = p_4 p_4 p_5 = p_5$

Substitution of the hydrogen atoms of anilms and its homologues an also be effected by and or chlorous groups, both in the hemone-nucleus and in the amido-group. In the latter case the derivatives are called an 11 do so (ngas 67), and are quite different from the bothes resulting from the substitution of chlorous randsals for the benzeue hydrogen. In the compounds of the latter class, the beare character is either much weakcased or entirely destroyed. Most of the deluvirties exist in several isomerom confections, seconding to the position of the substituting validates in the bearsen-nucleus. Examples of the locks of this class re-

And ine-sulphonic acid or sulphanilic acid, C₀H₄(SO₃H) NH₂
(page 49)

Natraniline, CoH4(NOg) NH2 (page 50)

Bromanilue, CaHaBi NH2

Trichloraniline, C.H.Cl. NH.

Mixed substitution-products, belonging at once to two or more of the foregoing classes, are obtainable by suitable means. As examples may be mentioned.—

Paramtracetanihde, $C_0H_4(NO_2) NH(C_0H_8O)$ Paramtroso-dimethylanihne, $C_0H_4(NO) N(CH_2)_0$

Paranitroso-dimethyl-paratoluidine, CoHg(CH2)(NO).N(CH2)a

The more important of the allies and derivatives of aniline formulated on this and the preceding pages are described in greater detail in the sequel.

On treating aniline, and also many of the above-mentioned homologues and derivatives, with oxidising agents, a series of brilliant colouring matters are obtained, which form the wellknown "aniline dyes" (Part I page 214 et seq.).

By the action of nitrous and, or a nitrite, on a cold solution of a sait of anime a sait of datas of a rea ne s obtained. Thus and the allied products obtained by similar means from the homologues and analogues of anime form the starking-point of the numerous and important colouring matters known as the "szo-dyes" (Part I. page 175 et seq)

In Diphenylamine and aniline hydrochloride cannot be caused to react with formation of triph on ylamine, $(C_0H_0)_0N$, but this body can be obtained by the action of mono-brombensone on dipolassium aniline — $2C_0H_0K_1+C_0H_0KN_2$, $C_0H_0K_1N_2$, $C_0H_0K_1N_2$, $C_0H_0K_1N_2$.

ANTLINE

43

By the action of reducing agents on the salts of diazobenzene, phenylhydrazine, $C_8H_8NH(NH_2)$, is obtained. The body has already been fully described (name 27)

Aniline.1 Amidobenzene Phenylamine,

$$C_6H_7N \,=\, C_6H_5\,NH_2 \,=\, \begin{array}{c} C_6H_5\\H\\H \end{array} \biggr\}\,N$$

Amilno occurs to a limited extent ready-formed in the products of the distillation of coal, hone, and peat of late years a small quantity has been actually recovered from coal-tar naphtha, but almost the whole of it is obtained inducedly from coal-tar by the action of a reducing agent on mitobenzene ("Amiline may also be obtained by passing aimnoma and benzene vapour through a relich titube $-C_{\rm H} + NH_{\rm g} = H_{\rm g} + C_{\rm gL}/N$. It is also formed together with diphenylations by the reaction of phenol and ammons. The best yield is obtained by heating phenol to alout 330° for twenty hours with ammonism chloride and magnessa or oxide of zme (oa ammonis-zme thloride, $Z_{\rm a}(NH_{\rm g}/C_{\rm b})$). Aniline is also obtained by numerous other reactions

And any be purified by fractional distillation and conversion into the acetyl-derivative. This is recrystallised from water, and on saponification yields pure aniline

Pure aniline is a colourless, oily hquid, of family vinous odour and aromatic, burning taste. It refracts light strongly, but has no rotatory action. Aniline, when very pure, freezes at 8° G, but a slight admixture greatly reduces its solidifying point. It boils at 183~184° Q, and distils unchanged.

The specific gravity of aniline is 10379 at 0° and 10216 at 20°, compared with water at 4°, and 10242 at 15°, compared with water at the same temperature. The coefficient of expansion is 000818

Auline becomes yellow or brown on exposure to air and light, especially at elevated temperatures, a resinous body being ulti-

Jantine was first obtained in 1836 by Un verd or be n by the dry dutilities on drinding, and necewid the name equitabless. Rung ein 1834 obtained it from ond-tax, and termed it squeed. The name cardiaes as due to Fi 1 to 0 is, who in 1840 dolanned it by distalling undape with castes alkabit. Then she kennadam was given it in 1842 by Zin in, who prepared it by reducing microbinesse by mighnisetted hydrogon. The names phesematic has also been proposed for it. Anthne was first accurately described in 1843 by A. W. Hoffm an in.

mately formed. The change is due to oxidation, and does not occur in vacuo or in the dark 1

Amine is only slightly soluble in water, requiring 31 parts at the ordinary temperature, but being more soluble in het water. Water also dissolves in amine, 5 parts being taken up by 100 of anitine at the ordinary temperature, and somewhat more at higher temperatures. The greater part can be separated by distillation, the water passing over first, but the last twees can only be iomoved by inclinated direction over cashe alkali.

Anilme is soluble in all proportions in a 50 per cent aqueous solution of its hydrochloide, and in smaller proportions in more dilute solutions (see page 67).

Aniline dissolves readily in alcohol, ether, wood-spirit, acatone, chloroform, carbon disulphide, and volatile hydrocarbons

Anilne is itself a solvent for sulphur, phosphorus, indigotin, camphor and colophony, but does not dissolve caoutchouc or copal. It is employed sometimes as a solvent for audine-blue

. Anilme is a powerful poison, coagulating albumin and producing symptoms similar to those caused by nitrobenzone (Vol II, page 478) ²

Amiline has marked base properties, a long sense of well-defined and crystallizable salte being obtained from it. It has, however, no action on phenol-phthalem, litnus or turnere, though it affects a few of the more delicate vegetable colours. It expels amounts from its salts at a bothing temperature, but is itself displaced in the cold. Amine decomposes the solutions of many metallic salts, with precipitation of the corresponding hydroxides. When heated with strong explainare said, and time is converted into para-amidoben 2 on e-sulphon ic a call of sulphamilic said. With hot framing sulphure acad, and it all phonic a call as produced.

¹ According to A Brdet (Compt. Rend., cvm: 520, Joser. Soc. Obers. Ind., vui. 883), authres and tolunduse propased by the reduction of pure nitro-derivatives are coloniloss after distillation, and though they become yellowed in a few days, light has us further effect on them, and even this change Bidel attributes to the presence of a mid oct-tin op hen. C, LHS. SNI.

* According to Lethe by and Turnbull the action of suttles a chiefly on the nervous system. According to Grand ho me, the first symptom in slight cases of possoning by anihus, caused by missing the vapor, is a bin collect on the edge of the large, while the gast become untstudy, the speech thick, the head effected, and the face pale, while the appetite fails completely accordingly. According agreement the symptom is In more series cases, and as any area from the saturation of the slothes with actions, the type become class bits or may be a superior of the contract of the children with a calling, the large because in the part of the contract of the contract of the children with a calling, the large to Wang to We had been all travel to be, much case as for example to the contract of the

In presence of an excess of acid, author imparts a deep yellow colour to pine-wood and alder-pith

According to Friswell, on adding cupric sulphate to an aqueous solution of aniline an apple-green crystalline precipitate is formed, or in extremely diluted solutions a green coloration

Cold aqueous solutions of amiline salts are converted by treatment with introus acid (or a nitrite and mineral acid) into salte of diazobenzene On boiling the solution phenolie formed, with evolution of mitrogen

Under the influence of oxidising agents aniline gives products and reactions which vary considerably according to the oxidiser employed, thus '—

- a When aniline is treated with excess of nitric acid, and the mixture evaporated at 100°C, the base is decomposed with formation of a brown substance. With smaller proportions of nitric acid various coloured modulets are formed, including pictic acid.
- b When treated with dilute eulphune and and manganese dioxide, aniline yields ammonia and quinone, CeH2O2, but the greater part of the product undergoes still further chauge.
- c If annune be diesolved in strong sulphinuc acid, and a few drops of a solution of potassium bichromate be added, a red colour is produced, which rapidly changes to deep blue
- d On treating amino, or one of its salts in a solid state, with strong sulphinic end, and then adding a minist fragment of menganese duxide or other oxidising agent (in the manner described under "strychinino"), a fine purple coloration is produced A better result is obtamble by employing electrolytic oxygen; in this form the test is the most deheste and satisfactory which can be applied.
- c Chlomme acts on dry annine with great violence, producing a black mass containing truch loraniine, Off, UC, M. Simme behaves similarly; and, on adding bromme-water to the aqueous solution of an amine suit, a precipitate of tribro multilute is formed. On the other hand, M. Ills and M. uter (Jone See Chem. Lnd, 1v 96) state that annino in solution in cabon disalphide reacts with Br., probably forming an additive compound.
- f When a solution of authne is treated with a dilute solution of bleaching powder, avoiding oxcess, a fine pupile coloration results, which gradually changes to brown When carefully applied, the reaction is delicate and characteristic after the colour is destroyed by other.
- q. If a minute quantity of aniline be treated with an aqueous solution of phenol, and a solution of bleaching powder be then gradually added, the reagent produces yellow strice, which change

to a greenish-blue. The test, which is due to Jacquemin, is said to be very delicate.

h If anilme, or one of its salts in the solid state, be testeds with a drop of cilicorform, and then solid potash or a strong solution of potash in alcohol be added, and the whole gently bested by immersing the tube in hole water, a specular blightly unpleasant odour will be produced, due to the formation of p hen yl -carban in c. CH_BNC The reaction, which known as "Ifofmann's isomitrile test," is produced by other aromatic monantum, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantumes, and by necessarily the produced by other aromatic monantum of the produced by the produce

DETECTION AND SEPARATION OF ANILINE

The foregoing colour-teactions are amply sufficient for the recognition of aniline, provided that a proper process of separation be previously amplied

Andline may be liberated from the squeous solutions of its salts by addition of caustic soda, and may then be extracted by sgriating the liquid with other On separating the othereal layer, and agitating it with dilute hydrochloric acid, the aniline passes into the aqueous liquid, which may then be concentrated or evaporated to dryness, and examined by the colour-reactions already described. From strychnine, which is the only substance with which aniline is at all ant to be confounded, it may be separated by adding caustic soda to the concentrated solution, and distilling over the anilms by driving in a current of steam, The strychnine remains in the flask, while the aniline will be found in the distillate if it be acidulated with hydrochloric acid and concentrated to a small bulk at 100° C. The same plan may be employed for detecting aniline in toxicological inquines, or the process used for isolating strychume may be used, but instead of evaporating the ether-chloroform it should be separated and agritated with dilute hydrochloric acid in the manner above described.

F Muller (Joss Ohem See, in 514) found unchanged annihm on the nume of a person possensed with it The turne was optically mactive, but reduced Fehing's solution. A portion of the concentrated urne, when boiled with strong hydrachloric and, neutralised with soda, and extracted with ether, gave an ethereal solution which showed the blue indophenol reaction. The othereal extract of the unboiled urne did not give this reaction, a fact which Muller believes was due to the secretion of the aniline as para-amidophenylaulphate (compare "Pinyl-Sulphure Acid," Part I page 9), a substance which is split by by bothing with hydrochloric and In support of this, the original turne contained sulphates (estimated by barnum chlorids)

equivalent to only 0 0475 gramme of sulphure and per line, A but after bothing with hydrobeliors each, 0 8085 gramme, A direct test for the presence of paramidophenyisulphates in unncounsist in boling the liquid with one-fourth of its measure of strong hydrochloric acid, adding a few cc. of a 3 per cent. solution of phenol, and then some drops of a chromic said solution. If para-simidophenol be present, the liquid becomes red, and turns blue on adding ammonia

The date mination of aniline may be effected by evaporating its eithereal solution, or preferably by extracting the best thereitom by agulation with dirtie hydrochlorie and, evaporating the acid liquid, and weighing the residual hydrochloride. Under favorasible circumstances it may be measured after liberation from a trong solution of the hydrochloride by addition of causic slight.

lustead of weighing the aniline hydrochlorids, the salt may be entisacted in water, and the solution titrated with standard silver intime Or it may be titrated with standard counts alkali and phenolphthalem or himns, as aniline hydrochloride acts on these indicators exactly like an equivalent quantity of fice hydrochloris and, and the end-reaction is perfectly sharp. The process allows of the titration of aniline in presence of neutral ammonicael salts On the other hand, with helianthin (meltiyl-orange), the basic character of free aniline is distinctly marked, but the cnd-reaction is not sufficiently definite to render the indicator available for accurately titrating aniline

According to Julius (Jour 800 Dyess, \$60, in. 79), freamline in aqueous solution can be satisfactorly titrated with standard sulphure or hydiochloric said, if congo-red be employed as an indicator and the neutral point be regulated as that at which a blumb-nolet colour is obtained, not changed by further small additions of acid, but a much largue access is required to produce a pure blue. Results are said to be obtainable agreeing within 0.2 per cent, with theory.

SALTS OF ANTLINE.

Annine combines readily with acide forming a series of salts which crystallise well. The following are the most important

Antine Hydrochlorade Hydrochlorate of Authne C₄H,N,HCl. The salt crystalbases with great faculty in colouriess needed of large plates, which are very soluble in water and alcohol. It lests at 192° C., and may be sublimed unchanged. It yelds double saits with stamme, mercure, antimomous, platime and sure chlorides, antime oblioryptatimate, (C₄H,N,HCl),FCl), crystallises from hot water in yellow needles. Antime sait is

the ordinary commercial name for anilme hydrochloride. It is manufactured by mixing the calculated weights of anilms and hydrochloric acid in stone-tanks, freeing the crystals formed from the mother-liquor by a centrifugal machine, and dying them. According to another process, anilms is dissolved in petroleum spirit of 0.720 specing gravity, and hydrochloric and gas passed in till the solution is asturated. The anilms salt is deposited as a white powder, which is separated from the adhening petroleum spirit by hydraulic pressure, and ground to nowder

Anthne salt is employed largely in calico-printing, its chief use being for the production of anilans-black (Part I page 250) It is important that the salt mended for this purpose should be made from pure amiline, and should be dry and neutral. The presence of free acid in the amiline salts is liable to cause the cloth dyed black to rot in the steaming process. It must be free from sand or gnt, which would injure the printing rollers, and will produce streaks on the printed cloth. Grit remains undissolved when the sample is treated with hot water, and may be filtered off, dried or ignited, and weighed Free acid is best determined by titration with decinormal caustic alkalı, usıng methyl-orange as an indicator, but the results are not very satisfactory A useful method of examination consists in titiating the aqueous solution of 2 grammes of the sample with normal caustic sods, using himus or phenolphthalein as an indicator The amount nontralised corresponds to the total acid, both free and combined with aniline Theoretically, 2 grammes of pure amline hydrochloride would require 154 cc. of normal caustic soda, but owing to the presence of toluidine and moisture commercial samples of good quality require between 14 and 15 c.c.1 The process will indicate the presence of ammonium chloride, which will not neutralise alkali, and hence a sample containing it will require a less volume of the standard solution Ammonium chloride is occasionally met with in considerable proportion as an adulterant of aniline salts. For its accurate determination the sample should be dissolved in water, excess of caustic sods added, the liberated amine senarated. and the aqueous solution distilled in the usual way. On titrating the distillate with standard acid and litmus or phenolphthalein, only the ammonia will be indicated Fixed impusities will be detected on igniting the sample; a mere trace should be present. An idea

^{.1} This method of examining aniline salts is due to R. Williams (Ohen. Netes, 1, 299), but he appears to attribute the reaction to the presence of free said.

of the proportion of tolaidine present in the sample can be obtained by hierating the mixed bases from the solution of the selfpeauly produced to the product of the selfpeaul quantity of strong areans easily collection to 180° of for some time. On boiling the product with water, the intensity of the crimson coloration will increase with the proportion of tolaiding in the sample. A more securate result can be obtained in the manner indicated on use 64.

Antime Sulphate, $(C_6H_7N)_2H_2SO_4$. This salt forms a crystalline powder, which is readily soluble in water and slightly so in alcohol. It is insoluble in ether, a fact which distinguishes it from the sulphate of methylamine

Antime Oxalate, (C6H7N)2H2C2O4, is very slightly soluble in cold water or alcohol, and insoluble in ether.

And the Acetate, C₆H₇N, HC₂H₃O₂, does not appear to have been obtained in a crystalline form When heated it loses the elements of water and forms a c et a n 111 de (see page 68)

ANILDER-SULFROND ACIDS AUDOSESCENE-SULFROND ACIDS When aniline is treated with an equivalent amount of dulue or concentrated sulphure and it is converted into suitine sulphate If an excess of acid be used, a high temperature employed, or sulphure anhydiade be present, antime-sulphone acid is produced—

$${\rm C_6H_5~NH_2\!+\!SO_2\!(OH)_2\!=\!C_6H_4\left\{\frac{SO_9{\rm OH}}{N{\rm H}_2}\!+\!{\rm H,OH}\right.}$$

Three modifications of this body exist, which differ according to the relative positions of the NH_2 and $\mathrm{SO}_2^{}\mathrm{H}$ groups in the benzene-chain Theortho-sulphonic acid (1 2) has no practical interest, but the meta- and para-acids are manufactured on a large scale for the production of anilinos—and aco-dyse

Meta-amidobenessessiphonic Acad, C,H,(NH,)¹⁰ SO,1¹², is employed for the neunfacture of metaula-g-yilon (Pat I page 190). It is prepared by warming intobenesse with funing sulphune acid, or by treating a solution of bensen on a stong sulphune and with funing nitric acid, when a mixture of n it ro-b en se n e sulphonic acid c, C,H,(NO,SO,H, is obtained, in which the meta-acid predominates, and may be roughly separated from its isomers by conversion into the barning or calcium sait. The metautive-sulphonic acid yields, on reduction, the corresponding amidosulbnone acid

Para-amadobenzenesalphona Acad, C₆H_ANH_N° SO, H°N, hkewase called Sulphamisc Acid, as prepared on a large scale by heating one part of aniline and three of concentrated sulphuric acid to 195°. With funung acid, the reaction occurs more rapidly and at a Vol. III PART II. lower temperature On pouring the cooled product into water, the acid separates as a crystalline mass, which can be recrystallised from hot water.

Sulphanile acid crystalliese in rhombic tables containing 1 agus, which affloresee in the air, and are only slightly soluble in cold, but readly in hot, water Theatment with potassium bichiomate and sulpharne acid exidese it to q i in on a Q₀H₀O₂. The solution of the sodium said, on irestiment with addium intits, yields so d in undia z obe nize ne sulphonate travels of all its base at 100°.

Audine subhamilate gives of flal its base at 100°.

NTRANILIESS When amine is treated with dilute intine coid it yields subme intrate. With the concentrated and it reneis far more violently than benzene, and is converted into quinon and other products. To obtain a inter-derivative by such means, the number must be protected by employing its acetyl-derivative, or by interfying im presence of excess of strong sulphume and to be batter case a mixture of the three isomers instainlines is obtained, but cheffy the medi-compound, in the former case paranitracet in 111 de, CHL(NO₂) NHC(CHLO), is formed, together with some of the ortho-compound, both of which readily yield corresponding nitra niline, CHL(NO₂) NH₂, on boing with concentrated hydrochions and or caustic fields.

Another method of preparing the intraulines, especially the mode-modification, is the reduction of the corresponding dunivobenzenes in alkalino alcoholic solution. Under these circumstances only one of the NO $_{\rm S}$ groups is reduced to NH $_{\rm S}$, whereas in send solutions of a mid ob enir en $_{\rm C}$, $_{\rm E}$, (NH $_{\rm A}$), is obtained (page 86)

	Nitra	TH ₂	
Appearance and	Ortho NOs NHs=1 2	Meta NOo NHo=1 3	Para NOs.NHs=1 4
Orystalline form,	Orange yellow needles	Long yellow needles	
Taste, .		Sweet, burning	Nearly tasteless
Melting point,	71*	114*	247°
Volatility, .	Distrils in a current of steam	Sublimes at 100° Distlis in a our- rout of steam,	Not volatile with
Salts,	Very unstable.	Bairly stable	Unstable
Behavious when boiled with strong soda,	-	Unchanged,	Forms pars mtro- phenol— C ₈ H ₄ (NO ₂) OH

The nutanilines are yellow crystalline bodies, readily soluble in alcohol but only slightly so in water. They are weak bases forming yellow salts, and yield the corresponding diamidobenzenes on reduction. The preceding table exhibits their chief differences

Two dustinantines, $C_0H_2(NO_2)$, NH_2 , are known, melting respectively at 182° or 138°. Also a trunt antine, $C_0H_2(NO_2)$, NH_2 , or pioramide, which melts at 186°, and is converted into pioric acid, $C_0H_2(NO_2)$, OH, and a m monia when boiled with caustic alkali.

Homologues of Aniline.

As already stated, the true homologues of aniline are bodies in which one or more atoms of the hydrogen of the betream-nucleas are replaced by a corresponding number of atoms of methyl or other alkyl radical. The compounds in question may be prepared, and are produced communically, by processes exactly similar to those which result in the formation of aniline. That is, the hydrocarbons tolucaes, xylence, &c., are treated with intrice acid, and the resultant nitro-derivatives are reduced to the bases by nascent hydrogen (usually iron and hydroboliton cacid).

In their general chemical relationships the homologues present the closest resemblance to annine, and yield substitution-products of a strictly parallel character They are also diazotised similarly

The only homologues of aniline which require separate description are the toluridines, C_0H_0N , and the xylidines, C_0H_1N . Then consideration will be followed by a section describing "aniline oils," under which term is included commencially pure aniline and tolundine, and various mixtures of these bases

Toluidings, Amidotoluenes Amido-methylbenzenes, Tolylamines,

$$C_7H_9N = C_7H_7NH_2 = C_6H_4(CH_3)$$
 H
 H
 N

The foliudines exist in small quantity together with aniline in coll-tar. They are produced commercially from toluene by processes exactly analogous to those by which aniline is prepared from benzen, and together with aniline constitute nearly the whole of the "aniline oils" of commerce (page 60). An interesting method of producing foliulune is mentioned on page 41.

Three isomeric modifications of toluidine are known. The chief physical differences between them are shown in the following table, in which they are also contrasted with aniline and their metamende be nzylamine, C.H.; CH., NH., 1

BENZYLAMINE is a colourless liquid of faint aromatic odour, and is not affected by light. It is miscible in all proportions with water, alcohol and

	Anshna	Ortho tolusione OH ₈ NH ₂ =1 2	Meta tolusdane CH ₃ NH ₂ =1 3	Para tolundins OH ₈ NH ₂ =1 4	Beneylamını
Specific gravity at 15°,	1 0288	1 0037	0 998 (at 25°)	Solid	090
Melting-point, .	Solidifies at -8° C	Does not soli dify at -20"	Does not solt- dify at -18°	Melts at+45*	Liquid.
Bolling point,	188* 7	199*	197*	198*	185°
Characters of the acetyl-deriva- tive —					
Melting point,	114*	107°	65°-66"	147*	57°-61°
Bolling point,	295°	206°	802°-804°	807°	800°
1000 parts of water dissolve,	8 4 at 14*	8 % parts at 19"	4 4 parts at 13*	0 89 at 22°	Soluble.
Solubility of the				()	
In 1000 parts of water at 15*,		28 8	26 5	87	
In 1000 parts of ether at 16°,		0.50	Very slight	0 016	

Ofthe-foludies is formed by the reduction of orthe-intertolucine. It is a colourless liquid, turning bown on exposure to air or light, and otherwise closely resembling aniline. It differs from its isomerides by griving a green coloration when treated with ferric chloride and a little para-diamidobenzene. A solution of 1 in 10,000 gives a fairly deep coloration, and one of 1 in 10,000 assumes a distinct greenist int. All commercial aniline gives this reaction, and even that prepared by the diskillation of miling with caustic alkali.

Meta-tolustine is produced by the reduction of meta-introducing, preferably by an and solution of stamous chloride. It is only present in small proportion in commercial toluidine. For its detection and approximate determination the mixed bases are convoxed into hydrochloridies, and the greater part of the someric salts removed by fractional crystallization. The mother-injuor is overporated to drynos, and the residue heated with unchyl alcohol to 200°, under pressure, for a considerable time. This produces a mixture of the three isomero dumsthyl-toludines,

other, but is separated from its aqueous solutions by caustic alkalies (compare "'Fyratine"). It has a strongly alkaline resolute, fumes with hydrochloric acid, and absorbe earloun dioxide from the air, with conversion into silky needles of the archands.

but only the meta-modification yaids a nitros-o-dorivaire, $C_{\rm eff}(NO)(C_{\rm H})_N$ and adding sodium interts to an escold solution of its hydrochloride. The hydrochloride of nitrosodium ethy lim eta-to-lui din o thus prepared, crystallises from a hot aciduated solution in greenish-yallow needles only slightly soluble in cold water On itsuatment with sodium carbonate the free base is obtained, melling at 92°, crystallising from water or ether in small green plates or long noodles, and prescription in moss-green needles on adding petroleum other to its chloriforms calution. All its solutions have a deep green colour Nitros-dimethylmetstoludine forms steel-blue compounds with annine and orthocloudine

According to Rosenstiehl, the three modifications of toluidine may be distinguished by the following reactions —

·	Os thotolussime	Metatolusdins	Paratoluidine
1 To a solution of the base in sulphuric acid, of 175 sp gr, add a solution of chrome acid in sulphuric acid of the same strength	Blue coloration changing on dilution to a permanent red- violot	Yellow brown coloration, be coming greenish- yellow on elight dilution, and colouriess on further addition of water	Yellow coloration
2 To a solution of the base in sul- pluric seld of 175 ap gr, add marie acid	Orange coloration, or in very con- centrated solu- tions, blown, be- coming yellow on dilution	At first ted, rapidly changing to in tense blood-red, and then dirty red, on dilution, orange	Biuo streaks which soon tinge the whole liquid, (in presence of ani- fine or ortho- toindidne, blood red) The colour quickly becomes violet, then red, and, after some hours, brown
2 Dissolve the base in either, and add an equal volume of water Then add not arons of clean solution of bleach ing powder	The aqueous layer becomes first yel low and them brown The ethereal layer, after separation, gives a per manent reddish violet coloration with dilute sulphurio acid	The aqueous layer becomes a thick buownish relieve The ethereal layer becomes toddish, and atter exparation and addition of dittle sulphurio acid is coloured violet at the under-surface	No reaction In presence of ant- line the other becomes blue on agitation.

Para-coloudne is produced by the reduction of the introduces derived from the toluens produced by the dry distillation of Toli balsam; also by heating paracresol to 300° with aminoma and chloride of zinc; and by molecular transposition from methylamline hydrochloride (page 41) It crystallases from hot

dilute alcohol in colourless plates melting at 45°, and has a peculiar odour recalling that of annine

Commoncial Toliudina consists chiefly of a mixture of the ortho- and para-modifications. According to Friswell, the specific gravity of the cithicolsuidina of commerce should be about 1-0037, and its boiling-point from 197' to 198' C. It ought not to solidity on cooling to -4', though the majority of samples contain sufficient paraticulation to cause them to commence crystallising at this temperature. The paratoliudina of commerce occus in white day crystals, suelts at 43'-446', and distalls between 196' and 198's. Laquid commercial toliudina should boil at 197'-198', have a specific gravity of cloud 100, and contain from 30 to 40 per cent of paratoliudine and 60 to 70 of orthotoliudina.

A portion of the para-modification separates from the commercial mixture of the isomers when the liquid is cooled by a fiscaning mixture. A further separation is effected in junctice by fractionally saturating the mixture of the bases with sulphilus soid, and then distilling in a current of steam. Orthochudine being a weaker base than the para-compound, the former will alone pass into the distilling if the quantity of sulphilus caid employed be somewhat in excess of that requisite to neutralise the paratoliuthes.

L Lewy (Jour Chem Soc, 1 872, Jour. Soc Chem Ind., v 481) has proposed to separate ortho- and para-toluidine by converting the bases into phosphates. It appears that when paratoluiding and oithophosphoric acid are brought together, di-toluiding orthophosphate, (C,H,N),H,PO, is produced as a salt ciystallising m scales and very sparingly soluble in cold water, but more readily, with partial dissociation, in boiling water. Aniline acts similarly, forming a sparingly soluble de-aniline orthophosphate, (CaH,N)2HaPO4 On the other hand, orthotolundine forms a mono-toluidine orthophosphate, (C,H,N)H,PO,, and never a dior tra-salt Hence in the phosphates obtained from a mixture of the two toluidines the proportions of the bases might be deduced from the percentage of phosphoric acid. The mono-orthotoluidine phosphate is more readily soluble in water than diparatoluidine or diamline phosphate. Further, when its solution is shaken with free aniline or paratoluidine, the orthotoluidine is set free. Hence pure orthotoluidine can be obtained from commercial toluidine 1 by adding rather more of a 21 per cent, aqueous solution of phosphoric acid than will suffice to form diphosphates

¹ The xylidines and cumidines behave like orthotoluidine, and form only monophosphates.

with the anilne and paratoludine present. On warming the hquid, the free orthotoludine forms a layer at the surface, which may be separated and distilled. The process may be modified by adding a further quantity of phosphate to convert the orthotoludine into monophosphate, and then cooling the hquid allowing it to stand to secure the complete deposition of the variatolidine intonephate.

Wolfing (Ber, xx 2182) states that orthofoludine prepared by Lewy hmaself by the above process, both on the and and large scale, stall contained as much as 4 per cent of paraticulume. For the preparation of pure paraticulume has recommends (Dungl Polyt Jour, celxin 200) that the hydrochlorides of the bases should be trasted with a muonit of south unitrite only sufficient to convert the orthofoludine present into a mid o a zo tol in en. Only when this change is complicadose the parateludine seast with the inhite to form a diagonuido-compound.

A method of determining the proportions of the ortho- and paramodifications of toluidine in the commercial product has been based. by Rusenstiehl on the different solubilities of the acid oxalates of the two bases The acid oxalate of paratoluidine requires 6660 parts of other for solution, while the corresponding salt of orthotoluidine dissolves in 200 parts of ether The method, somewhat modified, is as follows -0 2 gramme of the sample is dissolved in 80 cc of anhydrous ether free from alcohol. 1 059 gramme of anhydrous oxalic acid, or 1 177 gramme of the crystallised, acid is dissolved in 250 c.c of anhydrous, alcohol-free other Each c.c. of this solution will precipitate 0 005 gramme of toluidine An excess is added to the ethereal solution of the sample, the liquid allowed to stand in a stoppered bottle for twelve hours, then filtered through paper, and the precipitate washed with ether The precipitate is then washed into the bottle with water, and the solution intrated with decinormal caustic alkali and phenolphthalein. 1 cc of decinormal alkalı represents 0 00535 gramme of paratolundine Miniati, Booth, and Cohen (Jour Soc. Chem Ind. vi 419) find that if too long a time be allowed for the precipitation, the product is hable to contain the orthotoluidine oxalate, and hence the result will be above the truth. They recommend that a repetition of the experiment should be made, in which the amount of oxalic acid solution used is only that requisite to combine with the paratoluidine found by the first test, so reducing the erior to a minimum.

G. Lunge (Chemische Ind., viii. 74; Jour. Soc. Dyers, &c., 1. 150) estimates the proportion of para- and ortho-toluidine in a

mixture of the two by a careful observation of the specific gravity. The determination is made by the bottle, and referred to water at 15° C If the sample does not contain more than 50 per cent of paratoluidine it is liquid at 15°, and consequently the observation is made at that temperature With 50 to 60 per cent of paratolundine the method is still available if the bottle be filled at 20° C., but with still larger proportions the results are unreliable, as the correction for temperature loses in accuracy, and the differences in specific gravity become very small for considerable alterations in the composition of the mixture It is very desirable to adhere rigidly to the prescribed temperature, as an error of 1° C. causes an error of 7 per cent. in the estimation. The correction is ± 0 0008 for 1°, when the density is above 1 0008, and ± 0 0007 when below that point. All water must be removed by treating the sample with powdered caustic potash and redistilling. The distillation also serves to show the presence of analine or xylidine, in presence of notable quantities of which the method is inapplicable

Lunge gives the following table of densities of mixtures of para- and ortho-toluidine, water at 15° being taken as unity —

Specific	Ortho-	Specific	Ortho	Specific	Ortho	Specific	Ortho
gravity at	tolnidino	gravity at	toluidme	gravity at	toluidine	gravity at	tolundine
15° 0	Per cent	15° C	Per cent	20° C	Per cent	20 C	Per cent
1 0087 86 85 84 82 82 82 83 80 98 98 92 92 92 92 92 92 93 94 94 94 94 94 94 94 94 94 94 94 94 94	100 99 93 97 90 94 92 91 91 91 91 91 88 88 87 87 88 88 88 88 88 88 88 88 88	1 0018 15 14 13 12 11 10 08 08 07 06 06 06 07 07 06 08 08 01 1 0000 0 9090 98 97	824 82 81 80 784 77 76 75 74 72 71 70 69 684 684	0 9095 94 93 92 91 90 89 88 87 86 88 88 88 81 80 70 70 77 76 9 9075	061 85 64 62 61 61 60 66 65 66 66 64 65 65 66 66 66 66 66 66 66 66 66 66 66	0 9009 353 353 353 353 353 353 353 313 200 203 203 203 204 204 205 205 205 205 205 205 205 205 205 205	50 491 482 483 401 401 45 443 443 443 443 443 443 443 443 443

 toluidme has been based by G A Schoen (Chem Zeil., xii 494; Jour Soc Chem Ind , vis. 594) on the intensity of the red colour produced with potassium bichromate. If the specific gravity indicates the presence of more than 8 per cent, of paratoluidine it is reduced below that proportion by adding orthotoluidine 1 cc of the oil is then dissolved in 2 cc of hydrochloric acid and 30 of water, and 1 cc of a cold saturated solution of bichiomate of potassium added. The mixture is allowed to stand for an hour, with occasional stirring, and is then filtered Orthotoluidine gives a black lake and a colourless liquid, but in presence of para-. toluiding the precipitate is light brown, and the filtrate has a red colour, intense in proportion to the paratoluidine present Puie aniline behaves like orthotoluidine, but in presence of the latter a red filtrate is produced. Hence aniline must be absent, or its amount must be deduced from the boiling-point and specific gravity of the sample, and a corresponding amount added to the standard mixture with which the sample is compared

XVIIDINES Amido-dimethylbenzenes CoHg(CH)2 NH2

Six isomeric bodies of the above formula are theoretically possible, and all of them are known Thus 1—

	Positions of Groups CH ₈ CH ₅ NH ₂			Acetyl Derivative			
Base			Boiling- Point, " C	Molting Point, ° C	Appeniance, do	Characters of Hydrochloride	
s Orthoxylidine,	1	2	8	223	184	White needles	Moderately sol- uble white needles, con taining 1 sq
a-Orthoxylidine,	1	2	4	228 (molts at 49)	99	Long vitreous prisms	Long, very thin prisms, con- taining 1 aq
v-Metaxylidine,	1	3	2	214	176 8	White needles, not sapons fied by bool ing alkali or acid.	Thin anhydrous plates, readily soluble
a Metsxylldine,	1	8	4	219	129	White needles	Anhydrous rhom ble tablets, alightly sol- uble in cold water
s-Metaxylidine,	1	8	6	220	140 5	Large flat needles	Large anhydrous needles
Paraxylidine,	1	: 4	8	212-5	189	Long lustrous needles	Flat needles or large tablets

The table is chiefly drawn up from the descriptions of the isomene xylidines given by Roscoe and Schoilemmer (in part iv. page 406). The

The modifications of xylidine produced by nitrofying the xyler extense of coal-tar naphtha and reducing the nitro-derivatives are chief ant of α-orthoxylidine, α-metaxylidine, and paraxylidine, but two of the other isomere are also said to be produced. Only the a-metamodification is of any value for the manufacture of azo-colouring matters, and of the cumidines, CaHa(CHe), NH2, which are prepared by heating xylidine hydrochlorides with wood spirit On this account, the useless isomers are removed as far as possible from the metaxylene before nitrofying (Vol II page 483), and m fact the presence of even a few units per cent of orthoxylene will occasion considerable practical inconvenience by the formation of tarry matters during its conversion into xylidine On the other hand, commercial xylidine often contains as much as 25 per cent of paraxylidine v-metaxylidine (1 3 2) is prepared by converting commercial xylidine into the sulphate, which is allowed to crystallise, and the base liberated from the mother-houor by alkali The fraction distilling between 212° and 216° is heated with acetic anhydride The v-meta-acetxylidide formed is not acted on by boiling for several hours with four times its weight of dalute sulphurae acad containing 25 per cent of HoSOs, but its isomers are decomposed. On cooling, the unchanged acetyl-compound separates, and after recrystallisation from hot water melts at

characters differ conviderably from those attributed to the secures by Wroblewaky (Amalon, cvni 91) Nofting and Pick (Berchke, vxi 3150), however, consider that Wioblewsky's orthopyldine was simply impure s-metavyldine, and give the following table of characters of vylidine saits —

	v Orthoxylidine	as Orthoxylidine	v-Metaxylidine	Wroblewsky's so called Orthoxylidine
Hydrochloride,	+ 1 H ₂ 0	+ 1 H ₂ O	+# H ₂ O, needles	+ ½ H ₂ 0
Solubility in 100 of water at 18°C ,	11 2	Very soluble	9.2	Vory soluble
NITRATE, .	Anhydrous	Anhydrous	Auhydrous	Ankydrous
Solubility in 100 of water at 18° C ,	66	04	22	27
NORMAL SULPHATE,	Anhydrous	Anhydrous	Anhydrous	Auhydrous
Solubility m 109 of water at 18° C ,	14	5.6	Very soluble	
ACID SULPHATE,	Is not formed	under ordinary	+ 2½ H ₂ O	+ 23 H ₂ 0
Solubility in 100 of water at 18° C ,	eone	ALIGOR	62	Very soluble

XYLIDINES 59

176°8 C. On heating it for some time to 200° C, with three parts of sulphuric acid containing 70 per cent of H_2SO_4 , the sulphate of v-metaxylidine is formed. This salt differs from the sulphate of the isomeric xylidines in its very ready solubility in water

a-Orthosphiluse (1:2 · 4) as the only modification of xylidine which is solid at ordinary temperatures. By gradually evaponating its solution in petroleum ether, it is obtained in thick monoclimic prisms, but when rapidly deposited, or caused to solidity quickly, it forms transparent vitrous sablets. It meltes 4.5°, and also in alcohol and ether. Its squeous solutions are not coloured by blosching powder solution. The hydrochloria is readily solible in water, but so the property of the coloured by solution water, but only slightly in strong hydrochloris and, its aqueous solution impacts on misms gribor colour of he-wood

a-Metaphdase (1 3 4), or admary xylutine, is best obtained by converting commercial xylutine into the hydrochlorida crystallising the product from water. Both the hydrochlorida and hydrochlorida are only slightly soluble in cold water. The last tisces of impurity on he removed from metaxylutine by converting it into the activi-derivative, and iscircidizing this body home bournes till it has a melting-point of 129°. It is then decomposed by subburne and

Paraxylidine (1:4.2) has a specific gravity of 0 980 It is prepared by treating commercial xylidine with fuming sulphuric acid containing sufficient sulphuic anhydride to convert the bases into sulphonic acids. The mixture is heated to 100° for some time, allowed to cool, and the solid mass pressed under water to separate metaxylidine-sulphonic acid in the crystalline state, or the hot liquid is poured upon ice, when the metasulphomic acid, being with difficulty soluble in dilute sulphuric seid, crystallises out The mother-liquor is neutralised with chalk, filtered, precipitated with sodium carbonate, and again filtered. On concentrating the filtrate, the sodium salt of paraxvlidine-sulphonic acid separates in nacreous plates, which are washed with a little cold water to free them from traces of the readily soluble meta-sulphonate The salt yields paraxylidine on dry distillation with ammonium chloride, while the sodium ealt of metaxylidine-sulphonic acid chars under the same treatment. Paraxyhdine may also be obtained by nitrofying and reducing paraxylene, which may readily be prepared from commercial xviene (Vol II, page 483)

Cuminines. Amido-trimethylbenzenes. $C_0H_2(CH_2)_0$ NH_2 Various isomerides of this formula are known. The solid variety of commercial cumulane is made by heating xyhdine hydrochloride and methyl alcohol together under pressure, to about 300°. The bases are laborated and converted into intrates, and the infinculty soluble mirate of pseudocumdine separated from the mother-liquor. The base is again laborated and distilled The fraction passing over between 230° and 240° crystillies on cooling, and consists of am id-op-set of our mene—

It crystallases from hot water in long needles, and from alcohol in large prisms, melts at 68°, and boils at 234°-236° When converted into diazocumene it can be used for the preparation of azo-colours by reaction with nephthol-mono- and di-sulplionic acids.

ISODURIDEN Amido-tetramethylbenzene. C_sH(CH_s),NT_s. When the hydrochloride of pseudoctandue is heated with methyl alcohol to 300°, the hydrochloride of isodurdance is formed. The free base, which also occurs among the bye-products of the manufacture of pseudocumidne, is an ody huqud which bools at 50°-255° and solidifies on cooling to cystalts which mids 1 ts. 4°.

AMIDO-PENTAMETHYLEENZENS $C_0(CH_3)_2\,NH_2$ This base is obtained by heating dimethyl- α -pseudocumidine with methyl iodide. It forms large white needles, melting at 151° and boiling at 277°.

Aniline Oils.

The term "aniline oils" is applied commercially to all the different varieties of aniline manufactured on a large scale, equally whether the moduct in question consists of nearly pure aniline, of toluidine, or of a mixture of the two. The method of manufacturing the different varieties of aniline oil is substantially the same, the composition of the product depending on that of the hydrocarbon employed The details of the method of manufacture are, of course, subject to variation, but the following is an outline of the method pursued in a well-known and ine works :---Crude coal-tar naphtha is redistilled to a temperature of 170° C The product of the distillation, called "once-run naphtha," is treated with strong sulphuric acid (sp gr 1845) which removes the bases, hydrocarbons of the ethylene and crotonylene senes, and some of the higher homologues of benzene. A subsequent treatment with milk of hime or caustic soda climinates the phenols and other bodies of an acid character The purified oil is washed with water and redistilled to obtain "50/90 benzol," and this when fractionated with the acid of a dephlegmating column at once yields 99 per cent benzol, toluol, and solvent nanhtha (compare Vol. II. page 487) Solvent naphtha is now generally further treated for the isolation of xvlene, but the benzols and toluol are directly converted into the nitio-compounds by placing them an a vessel surrounded with cold water, and gradually running in a cold, previously made mixture, of 150 per cent by weight of nitric acid of 1 4 specific gravity with 200 per cent, of concentrated sulphuric acid. When the reaction is complete the mixture is allowed to stand, and the lower layer of acid is tapped off and concentrated again in glass for repeated use. The nitrobenzol is washed several times with caustic sode, and then treated with onen steam to drive off unchanged benzol and "light stuff" The nitrobenzol (or nitrotoluol obtained in a precisely similar manner) is then placed in a still with hydrochloric acid, and borings or filings of grey cast iron added gradually High-pressure steam is blown in, and the nitrobenzol which distils over is separated from the condensed water, and returned to the still until the complete solubility of the distilled oil in hydrochloric acid shows that the reaction is complete Milk of lime is then introduced, and the liberated aniline distilled off by the aid of steam Aniline sinks to the bottom of the condensed water, but when toluidine is being made the oil floats on the surface. The condensed water contains from 2 to 3 per cent. of dissolved bases, and is converted into steam for the aniline stills The iron is converted into a black paste, consisting chiefly of Fe.O., which is sold for purifying gas. The aniline oil is distilled to separate water, &c The addition of lime to liberate the aniline is not strictly necessary, and in many works it is omitted. The first reaction seems to be -

$C_8H_8NO_9 + Fe_9 + 6HCl = 3FeCl_9 + C_9H_8NH_9 + 2H_9O_9$

The ferrous chlorate formed also acts as a reducing agent, being converted into forme chlorate, which in presence of water gives ferric oxide and amine hydrochlorate. The end-products are chiefly amine, ferrose-ferric oxide, and a weak solution of ferrous chlorate. The hydrochlora and seems to act chiefly as a carriar, so that the general reaction may be imperented by the equation $-4O_{\rm th}^2 N_{\rm c}^2 + FF + 4H_{\rm s}^2 O = 5F_{\rm c} O_{\rm c} + 4O_{\rm c} \Pi_{\rm c} N_{\rm s} N_{\rm c} A$ scate acid was formerly employed in place of hydrochlora each, but its use is now simple, the control of the control of

COMPOSITION AND ASSAY OF ANILINE OILS

There are three leading kinds of aniline oil now recognised in the market, namely —(1) Pure aniline oil, (2) aniline oil for red, and (3) toluidine ' The demand for xylidine for the manufacture of axo-reds has considerably influenced the character of commercial anilms; since the 50/90 benzol, which was commonly used for the manufacture of *amiline for red," formedly contained a notable quantity of xylene, which is now removed and convexted separately. Since the employment of dephlegmating columns has become usual, benzone and tolenes of almest constant boiling-points have been manufactured. From the pure hydrocarbons the corresponding bases are prepared, while from the intermediate oil, containing about 25 per cent of benzene and 75 of tolsuens, an anilme oil for red is manufactured, which contains about 25 per cent of smilns, from 20 to 25 of paratoluidine, and 45 to 50 per cent of orthodoluidine.

In addition to the foregoing leading qualities of aniline oil, products of very varying composition and degrees of purity have to be dealt with by the dye-manufacturer Thus in making magenta by the arsenic acid process, fully one-fourth of the annline distils off and is condensed. But this recovered annline is found on rectification to have a considerably higher density than the original oil (1015 to 1009 against 10075), and to consist almost entirely of aniline and orthotoluidine, whereas the original oil contained from 15 to 25 per cent of paratoluidine. This is either employed for the manufacture of safranine or very jed shades of blue, or coude negatoloidine is added to it in such proportion as to bring it approximately to the original composition Similarly, in the manufacture of magenta by the nitrobenzenc process, the recovered aniline contains notable quantities of nitrobenzene, while from other processes methylated and ethylated andmes are obtained Recovered anilines are deener in colour and of greater body than unused oils, and often have a strong and somewhat characteristic odour They are rarely met with outside the colour-works in which they have their origin.

On next page 1s a tabulated last of the more unportant or frequently-cocurring constituents of antline oils 2 With the exception of aniline and its homologues, and the substituted milines, very little 1s known respecting the effect of the bothes formulated in the table on the colouring matters produced. For the most part the objectionable impurities are got rid by fractionating the crude aniline oil

¹ The composition of amine oil for red is often judged of by the consumer solely from the specific gravity, and he or the amino-makes adjusts it accordingly by adding amine or tolundine to the crude oil as the gravity may makeste

² Hell and Rockenbach (Ber, xm 505) have investigated some other non-basic constituents of author and tolundue tailings

Name	Formula	Melting- Point "U	Boiling Point C	Remarks
Amiline,	CeHa NHa	- 8	188 7	See page 48
roim (artho-, 1 2 meta-, 1 8 para-, 1 4	CoH (CH2) NH3	below - 20 below - 13 45		See page 52
Xylidine (several isomers), Jumidine (several	C ₆ H ₈ (CH ₆) ₂ NH ₂		212-226	See pagé 67
isomers, chiefly Pseudosumidine),	C ₆ H ₂ (CH ₃) ₃ NH ₃	63	285	See page 60
Methyl anilme.	CaHa NH(CHa)		192	See page 73
Dimethyl-aniline	CaH3 N(CH3)2	0.5	192	See page 74
Ethyl-antline.	CaHa NH(CaHa)		234	See page 73
Dinhenylamine.	CaHa NH(CaUa)	54	802	See page 79
Acetanilide, .	C_0H_0 NH(C_2H_0O)	118	295	See page 68
	C*H*(CH*) NH(C*H*0)	65-86 147	802-304 900-807	(Produced by action of heat on tolukline
Nitranilines,	C ₅ H ₄ (NO ₂) NH ₂			From imperiost reduc- tion of dimiroben-
Paraniliue.	C12H14N2	192	330	
Xenylamme.	C12H9 NH2	45	322	
Phonylene diamine (para),	C6H4 (NH2)2	63	287	Reduction of dinitro- benzene (page 87)
Toluylene diamine (para).	CaH2(CH2) (NH2)2	99	253-255	See jinge 88
Azobeuzene.	CaHa Na CaHa	65	298	Imperfect reduction
Nitrobenzene,	C ₆ H ₈ (NO ₂)	118	210	of nitrobergene Vol II page 478 Monoglinio tables
Dinitio meta	ChH4(NO3)2	90		Long needles or thir
Nitro para ortho meta-	C*H*(GH*)(NO*)	172 below - 2 16 51	928 200 200	Monoclinic needles Sp gr 1 168 at 23' 5 Sp gr 1 168 at 23'.
Benzene,	C _d H ₆	5.5	80 8	Vol II pago 469
Toluene,	CoHe(CHa)	below - 2	111	Vol II page 479 ,
Amidothrophene,	C4H2S NH2	1	1	
Paraffins,	C ₂ H _{2n+2}			Repecially in anilone offs derived from cauncil tar housels

The assay of anilue oils is usually lumited to observations of the colour, odour, and specific gravity, supplemented by a careful fractional distillation and tests for water, nitrobenzene, hydrocarbons, &c.

The specific gravity of annine oil is a valuable indication of its composition. The observation must be made by the plummet or specific-gravity bottle at exactly 15°C., and the result referred to water at the same temperature taken as unity 1

¹ P Schoop (Chem. Zett, xx 178, Jour Soc Chem. Ind., v 178) gives the density of pure aniline as 1 0877 at 1°C, orthotolundine as 1 0143, and partolundine as 1 0045 at the same temperature, the coefficient of expansion being in each case 0 00081 for 1°C. The following figures represent the densities as thus observed -

	Specific gravity at 15° C
Pure anilino, .	. 10268
Aniline oil for red, .	. 1 0075 to 1 0012.
Orthotoluidine, .	. 1.0037.
Mixture of equal parts of and para-toluidine,	ortho- } 9975.
Do and along James	. Solid.

The odore of pure and the is very different from that of the toludues II he presence of bloudues in annine is indicated the density of the sample, its dimmished solubility in didta slockol (rage 68), and by the results of the factional distillation, define 66) In addition to these characters, the following tests are sometimes of sorries—

Pure ambine affords no resembine on treatment with excitating agents, but if cloudne be present magents is readily formed. The test is best made by mixing 5 c of the sample of ambine with an equal measure of a concentrated solution of arsence and, containing about 75 per cent. of As₂O₅ and having a density of 2 04. The mixture, contained in a small flask or long test-tube, is immersed in a parafin-bath heated to 180° C. The mixture rapidly changes in coloni, and swells considerably. When the action is complete, the contents of the tube acquire a metallic browse appearance and no longer intumesee. The product is treated with boiling water, when, if the sample contained follution, extended to the high continuous considerable containing dissolves and communicates an intense ermson colour to the liquid. Neither pure antime nor follution along gives this section.

If a sample of commercial anilane be mixed with some solid magenias and a few drops of glaceal aceae acul, and the whole heated to 180° C, as described above, ammonia is abundantly evolved, and in a short time the mixture bocomes intensely blue from the formation of triphenyl-rosaniline. With pure anilane his blue is very pour on shade, but when tolender or xylidine is treated in a similar manner the product is intensely purple, and a mixture of the bases gives proportiousts intensely purple, and a mixture of the bases gives proportiousts intenselying the shades of colou. If a little of the "mott" be withdrawn from the tube, diluted considerably with alcohol, a few drops of acetic acid added, and then strucked on white filler-pere by uncan of a glass rod, the purple tim is readily observed, especially if the paper be held up before a gas-flame

A valuable indication of the general composition of an amiline oil is obtained by submitting the sample to fractional distillation, and noting the proportions of distillate obtained at various tempenatures The distillate may be measured after each ruse of 5 dagrees in the boling-point of the sample, or the temperature may be observed when each consecutive 5 or 10 per cent fraction has passed over The latter is the plan now commonly adopted, 100 or of the sample being employed, and the ariangement of the apparatus being exactly the same as in the fractional distillation of bonoils (VGI LI page 495).

The heat is applied cautiously at first, in order to disspate any water. When thus is effected, which will be known by the rand rise of the thermometer, the heat is so regulated that the distillate shall fall in distinct drops, about early per minute. With each norcease of 10 cc. in the volume of the distillate the temperature industed by the thermometer is observed and recorded, the process being continued till 90 or 95 cc. have passed over

A vay simple test for aniline oils was dovised and communeated to the writer by the late B Nickels, who found it to give useful results, and to indicate differences between samples not readily distinguishable by the ordinary fractional distillation process. The test is based on the greater solubility in diute alcohol of aniline as compared with toliudine and xydinies, and is thus performed—5 c.e. measure of the sample is taken with a pipette and diuted to 40 oc with methylated spirit. Distilled water is then gradually added from a burstle, with constant shaking, till a permanent furbility is produced, when the volume of water employed is noted Operating in this way, a sample of very pure aniline required 126 c. of water to produce permanent turbidity. The following figure, obtained by B Nickels in 1881, show the results yielded by three-virgual specimens of commercial aniline as the namifactured.

	Pure Antilne	B Heavy Anilline	Toluidine.
Colour, Specific gravity at 15° 5 C Water required for precipitation,	Pale amber 1 025 106 4 c c	Amber 1 011 78 7 c c	Deep brown 1 992 63 2 0 0
10 per cent distilled over at . 20	* C 1831 1831 1831 1841 1841 184 184 184 184 184	189 1895 1800 191 1913 1922 189 1943 197 291	195 1952 196 198 198 197 197 197 198 199 199 199

Sample A was a fair commercial specimen of the quality known as "pure aniline," and actually contained some 95 per cent VOL III PART II

of real annine. An article of this high purity is required for the namufacture of annine blue, triphenyl-rosaniline (see page 64), any notable admixture of tolundine resulting in a product dyeing with reddish tinge 1

The quality known as "heavy anline," exemplified by B, as fair sample of anline oil for red (see page 83). The class of annihne is produced from benzols containing a considerable proportion of toluene, and the annihne oil trailer is a mixture of annihne and tolumines. Good samples of annihne oil forted contain from 35 to 42 per cent of real annihne, 35 to 50 per cent, of particularities.

R J Friswoll thinks 100 cc an undesirably small quantity for fractional distillation. He prefers to operate on 250 cc, which he distills in a flack with a side-tubulure, and he recommends an observation of the temperature at which the last drop disappears from the bottom of the flack. A naked flame is used, and a few fragments of platinum wire or fire-brick added to the contents of the flack. The following figures were obtained by Frawell (Thorpe's Dett Applied Ohem, 1, 165) by the examination of commercially pure aniline.

	No t	No 2	No 3
Specific gravity at 15° C,	1 02710	1 02684	1 02690
	*c	• 0	.0
10 per cent over at, 20 11 12 12 13 14 14 14 15 15 16 16 17 17 18 18 18 18 18 18 18 18 18 18 18 18 18	184 7	184 6	184 6
20	184 7	184 8	1816
80	184 7	184 8	184 7
40	1847	184.8	1847
50	1848	184.8	184.8
80 11 11	1819	1848	184.8
70	185 0	181 8	194.9
80 0 0	185 1	1818	184.9
90 , ,	186 1	184 8	185 0
" Dry at,	1867	180 8	

Any scate present in aniline oil will be found in the very first protrions (first fraction of 10 per cent) whenever the sample is submitted to distillation. It takes the form of globules, which are not miscible with the next fraction of the distillate nor with petroleum spirit. Water may exist in maline in any proportion from a trace up to 8 or 4 per cent, but a good commercial rectified specimen should not contain more than 0.5 per cent. Annine is readily soluble in a strong aqueous solution of aniline hydro-

¹ In good samples the boiling-points hold closely together, differing by one or two degrees only Inequalities or jumps in the boiling-point, especially at the beginning and end of the distillation, indicate badly-in-ide samples or mixtures.

chloride A solution of the kind, of 108 specific gravity, is stated by Watson Smith to be sometimes sold as annine oil, which in colour and taste it closely resembles. Such a fraud would be at once detected on distillation

Benneau, tolutena, and othen hydrocarbons will separate when the first fraction of 10 per cent (10 eo) as treated with an equal volume or slight excess of hydrochloric and, and water added to 100 or 150 eo. They assume the form of only globules which float even and indiving the hquid. The best samples of pure anilme show only a slight opalescence when thus treated, but the smell of the "light staff" (Vol II page 485) as laways preceptible. In recovered anilmes these impurities exist to a notable extent, since they survive the reactions by which the bases are consumed. Amilne for red usually contains somewhat more hydrocarbons than pure anilme.

Mit obsesses and survolviese may be recognised, even when mere timese an epissent, by the milky appearance of the liquid produced by saturating 10 co of the original sample of oil with hydiochlore and On disting the high with wate, and leaving it at rest for some hours, any considerable quantity of nitrobenzene will collect at the bottom in the form of oily globules, which, aftes separating the acid liquid, may be identified by the smell and other characters. Still smaller quantities of mitrobenzene may be recognised if the "tailings" be operated upon, instead of the original sample Nitrobenzene occurs more frequently in magenta-aniline and toludite than in the oils of lower bouling-point.

Nitrobenzene is also indicated by the yellow colour of the fioth produced when the sample is violently agritated.

Acctantitide and accetoclutude were impurities characteristic of aniline prepared by the reduction of nitrobenzene with aceta caid and iron, but are now rarely met with in aniline oils. In any case they would become concentrated in the "tailings," together with phenyleno-dismuns, acceleracies primitine, "xenylamins," for

Anilms tailings us the name applied to the least volatile portion of anilm oils. They contain little or no anilms; some toludine, xylidine and cumdine, nitrobenzene and its homologues; and some or all of the bye-products tabulated on page 63 which but above 200°

The composition and special methods of examination of commercial tolundries are described on page 54 et seq.

Anilides.

The antildes are derivatives of aniline in which one or both of the hydrogen-atoms of the amido-group are replaced by acid-

radicals The homologues of annine yield similar derivatives (e.g., a ceto-tolulde, page 52). The most important and typical member of the class is a cetanilide or phenylacetanide — $C_0H_{\pi} \operatorname{NH}(C_2H_{\pi}O)$, or $C_3H_{\pi}O \operatorname{NH}(C_0H_{\pi})$.

A number of derivatives of acetainlide have been prepared, and certain of them have found some employment as analysesics and antipyretics, as for instance —

Acetanlide Phenylaostanule Antifebran $C_0H_3 NH(C_2H_9O)$. Bronacetanlide Antiseptin Brom- $C_0HB_1 N_1H(C_2H_9O)$. Methylacetanlide, Exalgin Methylacetanlide, Exalgin Methylacetanlide, Exalgin Methylacetanlide, Exalgin Methylacetanlide, Methocombeno H_3 or H_3 or H

Most of these bodies are described in the following pages The relationship of antifebrin to hypnone, hydracetin (pyrodine), and phenyl-arethane, is shown by the following formula —

Aestophenome Hypnone (Part I page 23) $C_{\rm eH_2}$ (CO CH₂). Acetanlude Antidebru (see below) $C_{\rm eH_2}$ $C_{\rm eH_2}$

ACETANILIDE PHENYLAGETAMIDE CAH, NH(CoH,O).

This substance was originally obtained by the action of acetylchlorido on anilme. It is more conveniently prepared by bohing amiline with glacial acetic acid for many hours under an inverted condience, until the product solidises on acoloning. The mass is then melted and poured into water, for remove unconverted anilms and acetic acid. It may be purified by distillation and crystallisation from alcohol, bensene, or hot water, from which it separates in colourless unctaous laminaes, resembling born acid, soliuble in about 190 parts of cold or 18 of boiling water. Acetamilde in about 190 parts of cold or 18 of boiling water Acetamilde in colouless, but produces a sight buruing sensation on the tongue It occurs commercially as a crystalline powder or scales. It mails at 112"-113", and distals unchanged at 295" C. Acetamilde in dissolves in $3\frac{1}{4}$ parts of alcohol, and is very soluble in ether, chloroform, and benzeue, yielding neutral solutions

Asstanthde is a weak base. The hydrochlorude is obtained by passing hydrochlorus acid gas through a solution of acetamhde in acetone. It forms needless which are decomposed into their constituents by water, and gradually converted into acetic acid and annine hydrochlorude on exposure to most air.

Acetanlide dissolves in strong subplume and without change of colour. On treating the solution with nitre acid, the sociatinate is converted chiefly into pare-nitro acid in little (page 50); some of the archico-compound and, in presence of a large excess of sulphume send, a little of the medi-compound being also formed Nitrons acid, passed into its acetic send solution, converts sectamilide into an unstable nitro as mine, CH₂N(C,H₂O)(NO). When hested with am oblicate to about 250°, acetamilde yaelist faccountiers, C₂H₂N(C,H₂O)(NI). Therefore in alcoholic solution with sodium ethylate, acetamiled yaelis as od it in derivative, CH₂N(C,H₂N)(H₂O), but when thus is boiled with water it splits into a nil line and sodium a cet at the Acetamiled behaves like amiliae on treatment with caustic alkali and chloroform (page 46), and the formation of the disagreeably smelling isomirile is a delicate reaction for its presence (compace page 83)

asonitrie is a denouse reaction for its presence (compare page 85)

Acetamilde behaves like aniline when treated with phenol and solution of bloaching powder (page 45)

When treated with a solution of potassium chlorate in strong sulphure send, acetamine gives a red coloration, changed to yellow on dulution. With a crystal of a mitrite and a drop of concentrated hydrochloric send it produces a yellow colour, changing on heating to green end blue; and, on evaporating the hund to dryness, an orange residue is obtained, changed to red, on adding ammonia (Yitali).

When noctamilide is heated gently with mercurous nitrate, a body is produced which dissolves in alcohol with green colour (Y v on). If a few centigrammes of acestanide be gently heated with two or three drops of a solution of mercurous nitrate, and when solution has been effected two or three drops of sulphurue and added, a blood-red colonation will be produced (Cella and Arren o). The same reaction is produced by phenol, resorund, thymol, and salecylie, gallie, and tannic acada, but not by benzouc and

Acetanilide gives no colour-reactions with ferric chloride, nitrites in very dilute solutions, or potassium bichromate in aqueous solution. These reactions distinguish it from antipyrine and kairine

Various other colour-reactions of acetamilde have been described

As a rule, the most satisfactory method for its positive identification is to heat the substance with alcoholic potash, dulute with water, and shake with ether. The ethersal layer is examined for amiline, while the aqueous liquid is tested for an accetate.

To detect acctanilide in urine, Vulpius boils the liquid with hydrochloric acid, cools, extracts with other, and tests the othereal solution with phenol and bleaching powder solution

E. Ritsert (Pharm Zeit, xxxv. 306, Jour Chem. Soc. lviii 1349) gives the following tests for the purity of commercial acetamilide .- The sample should leave no ash on ignition, and after diving for two hours at 105°, should melt at 114°. A higher or lower multing-point indicates the presence of aceto-toluides 01 gramme dissolves in 1 cc of strong hydrochloric acid to a clear solution, which, after a few minutes, precipitates acetanilide hydrochloride (methyl-acetanılıde docs not yıeld a sımılar reaction) No change should be produced on adding a drop of mirre acid, which, after a time, produces a yellow or brown coloration if phenacetin or methacetin be present. It 0.1 gramme be boiled in portions in 2 cc of strong hydrochloric acid, the solution cooled, and a drop or two of chlorine water added, a fine blue coloration is produced The aqueous solution of acetanulide should be free from acid reaction (indicating acetic soid). On boiling it and adding ferrie chloride, a deep reddish-brown colour should be produced. destroyed by a mineral acid. If a drop of dilute solution of potassium permanganate (1 1000) be added to a boiling aqueous solution of 1 gramme of acetanilide in 30 cc of water, the pink coloration at first produced should persist at least five minutes, and should not change to yellow on again boiling Procepitation at this stago indicates the presence of free aniline, resinous products,

In the additions (1890) to the Britch Pharmacoguezia, nectamidia is described as melting at 238° F (= 112° 8° C), and dasolving in sulphure and without coloration. The solution in I8 parts of bouling water should be clear, neutral, and odourless; and after cooling should not be coloused on adding ferre chloride this is directly opposed to the expenses of Ritsert above quoted. In the German Pharmacogueza the direction is to add ferre chloride to a cold astinuted solution, thus avoiding the dissociation and formation of aceta cand hable to occur on bohing. According to the German Pharmacogueza, no heating with caustic alkela solution, acetamide gives off an acomatic vapour, which, after addition of a drop of chloridorm and renewed application of heat; is changed to the disagreeable smell of the isomitrile. Further, O'I gramme of extending should yield a clear solution when bouled with 1 c c of sectaming should yield a clear solution when bouled with 1 c c

aceto-toluides, or other impurities

hydrochlone acid for one minute, and, after adding to the liquid 2 cc. of carboine acid, a cloudy red coloration should be produced by solution of bleaching powder, changed to a permanent indigobles (in do v be no 1) on adding excess of ammonia

Actanishe has powerful anthyrests properties, and has sconved an extensive application in modiume under the name of "an itife brin," though dangerous symptoms are sometimes produced by it (Plarm Jour., [3], xx 1058) The dose is from 3 to 10 grams. According to Salzer, commercial satisfains its lable to certain unchanged amiline, which may be detected by dissolving the sample in cold hydrochlore and, and pouring on the higud a solution of bleaching powder. Pure acctanished yields a white prequisite, which dissolves on shaking the liquid, but after a time colouries silky needles separate. In presence of amiline the well-known volet coloution is produced.

Acetaminde has been used as an adultment of antipyime (page 36). The melting-points of the pure substances are nearly identical, but a mixture of equal proportions of the two melts at 45°C Of the three isomeric aceto-tokudes (page 52), only the meta-

compound possesses antipyretic properties

Pana-hom-acetawitate, C₂H₂B.NH(CO CH₃), has been introduced as a remedy under the name of "a n tr s e ps in". It forms small pently prisms, melting at 164°5, and devoid of taste or smell It is soluble with difficulty in cold, but readily in hot water, as also in alcohol and ether

Accumentalylamitude on Mcthyl-accumulate, C.H. N(CH.)(C.H.C.), as prepared by warming together methylamiline and acetyl chloride. The product is boiled with water, when the new body crystallises on cooling. Methylacotamilide has been introduced as an anti-neumatic and analgeses under the name of "c x at g in " In dosses

¹ When administrated to rubbits, acctanable as conduct to pas-amulciplismos, CAL(LOST) NN₁, with complete demnation of the sectyl-group. In dege there is a small founction of pasi-amulciplescol, but the clust change consists in a smultaneous catation of the animin-residuate to ortho-amulciplismos), of the sectyl-group to carboxyl, and in the formation of carboxyl-ortho-physics of carboxyl and in the formation of carboxyl-ortho-physics of the complete of the complete

is excreted in the union as a sulphate. In both it he rabbit and the dog the smide phenois are also chimated as sulphates. In man, the acetyl-group is not whelly consider, the turne containing the sulphate of aceto-par am id op hen ol. In all cases there is an oxidation of one of the hydrogram of the hearness of the hearness—nuclous to hydroxy; while the proportion of edecad sulphate is inonessed (compare "Aniluse," page 46), the union is reflected sulphate is inoressed (compare "Aniluse," page 46), the union is reflected in the control of the control of

of $\frac{1}{2}$ to 4 grains its effects are said to be very satisfactory Exalgin forms fine needles or large white tablets (compare "Acctanlide") If melts at 100°-101°, boils without decomposition between 240° and 250°, and is slightly soluble in cold water, but moie so in boiling, and very soluble in water containing a little alcohol. It is saponified with difficulty by caustic alkali, but completely by concentrated hydrochloric acid, with formation of acetic acid and

Hirschaohn states that exalgin may be distinguished from antifebrn and phenacotin by treating I gramme with 2 c.c. of chloroform, which dissolves the exalgin only A chloroformic solution of exalgin remains clear on adding ten measures of petroleum ether, whereas the solutions of antifebrm and phenacetin become turbid 20 per cent of acetamhde, or 10 of phenacetin, may be detected in exalgin by these reactions An aqueous solution of antifebili gives a bromodelivative on adding bromne-water,

BENZANLIDE, CoHo NH(CO CoHo), is obtained by the action of benzoyl chlorde on anilme, or by boiling together equivalent quantities of benzore seed and amiline. It forms a white, crystalline powder, melting at 160°-161° and volatile without decomposition Li is almost insoluble in water, but dissolves in fifty-eight parts of cold, or seven of boiling, alcohol, anystallising on cooling in nacreous plates It is difficultly soluble in other. Benzanihde is not attacked by aqueous alkalies or acids, but is saponified by fusion with caustic potash It has been found valuable as an antipyretic for children, in doses of 2 to 8 grains, and is said not to produce objectionable secondary effects.

PHENTL-DEFTHANE ETHIL CARRANHATE C_0H_0 NH(CO OC_0H_0) This compound has recently acquired a practical interest owing to its introduction as a synthetic remedy under the name of "euphorin" It is produced by the reaction of anilme on ethylchlorocar bonate, and occurs as a white crystalline powder, of a family aromatic odour and scatcely perceptible taste, which subsequently becomes acrid and clove-like It melts at 49° to 51°, boils at 287°, and is only slightly soluble in cold water, but very freely soluble in alcohol, and sufficiently soluble in sherry and other alcoholte figures to be conveniently given in solution in such menstran

1 Exalgen may also be distinguished from antifebrin, methecetin, and phensectin by treating 2 grains (or 0.1 grainine) with 20 minims (or 1.0 c.) of actent by treating a gramm (or v. gramme) went as mining (or v. c.) or concentrated hydrochlorio acid. Phenacetin remains insoluble Antidorin dissolves, but separates again in crystals of the hydrochloude Methoscin also dissolves, but is recognised by the reddish-brown coloration produced on

to Sanson1, after administration of phonyl-urethane, the uruse shows the para-amilophenol reaction either directly or after distillation with poissium carbonate. The proportion of urea is increased, but the uruse is free from phenol, aniline, albumin, and sugar.

Substituted or Alkylated Anilines.

These bases result from the replacement of one or both of the hydrogen atoms of the amido-group of annine by alkyl or other basylous radicals.

The bases of this class are obtained by heating the hydrochloride or other salt of aniline (or its homologues) with the alcohol with which it is intended to react, or the halogen salt of this alcohol with free amiline.

The only substituted anilines which require special description are the following ---

	Pormula,	Specific Granty	Boiling Point	Reference.
Methyl-anilina, Dinethyl-anilina, Ethyl-anilina, Ethyl-anilina, Diethyl-anilina, Phenyl anilina (Diphenylamine), Diphenyl anilina (Triphenylamine),	C ₆ H ₅ NH(CH ₆) C ₆ H ₅ N(CH ₂) ₅ C ₆ H ₅ NH(C ₂ H ₆) ₅ C ₆ H ₆ N(C ₂ H ₆) ₅ C ₆ H ₆ NH(C ₆ H ₆) ₅ C ₆ H ₅ N(C ₆ H ₆) ₃	976 at 15" 9663 at 15" 964 at 18" 987 at 13"	192 192 294 213 5 202	Page 78 Page 74 Page 79 Page 79 Page 89

Diphenylamine is a very weak base, and in triphenylamine the basic character is entirely lost

METHYL-ANILINE. C.H. NH(CH.)

This base is obtained by the action of iodide, nitrate, or chlorade of methyl on aniline, or by heating methyl alcohol with unline hydrochloride I in all cases dimethyl-aniline is formed simultaneously, and hence in the production of mono-methylaniline a portion of the aniline remains. in uracino. unattacked.³

¹ Pure methylaniline may be obtained by the reaction of methyl indide on sodium acetanilide, C₂H₂ NN_N(C₂H₂O), and asponification of the resultant compound by causto alkali.

To squase that from its mone- and do motify-derivative, disite sulpture and is added to long as unline sulphate cortinues to spenter. The sulphure and solution is separated from the solid anime sulphate by pressure in a linear cloth, and the expressed luquid treated with camen soul. The substance which separates in dried and treated with eacily chloride until no further unit of temperature as cleared, when the product is poured into cold with Coccoling, me thyl-section 114, Q.H. NOCH_NOCH_SO, separates in long cases, which drum thyl is not in hydroci lord remnans in solution meetles, while drum thyl is not in hydroci lord remnans in solution.

Methylamine as a lagud bosing at 192°. It resembles anilus, but as lighter than water, and its odour as stonger and more aromatic. The sulpinet is soluble in either and uncrystallisable. As colution of bleaching powher fists colours it voiced and bleaching powher fists dolours it voiced and believe to the brown. The conversion of methylamine into toluidine is referred to on name 41.

Methylaniline-nitrosamine, CaH, N(CH2)(NO), separates as a yellow oil on treating a cold solution of methylamline hydrochloride with sodium nitrite, while any aniline and dimethylamiline are converted into soluble products. If the nitrosamine be extracted by ether, and treated with tin and hydrochlone acid, it is reduced to methylaniline, which may thus be obtained in a pure state (compare page 7) The mitrosamine is destitute of basic properties. It has an aromatic odour, and may be distilled in a current of steam, but not alone When methylaniline-nitrosamine is warmed with phenol and sulphuric acid, the nuxture diluted with water and saturated with caustic alkalı, it vields the intense gicen-blue coloration moduced by all nitrosammes (Liebermann's reaction) When heated with alcoholic hydrochloric acid it undergoes molecular transformation into paranitioso-methylaniline, CaHa(NO) NH(CHa), a body crystallises in green-plates or steel-blue prisms, and otherwise resembling paramiroso-dimethylamline (page 75) DIMETHYL-ANILINE CaH, N(CH2),

This impostant base is obtained by the action of excess of methyl nobile on authon. On the large scale, methyl icolds was formely employed, but was afterwards replaced by the minute, and this again (owing to its explosure proposteries) was superiseded by the city volatile methyl chloride. The product obtained in this way contained about 5 per cent of monomethyl-emiline, but no other adunxtures. Dimethylaulime is now always manufactured by leating together a mixture of amiline hydrochloride, aniline, and methyl alcohol? The methyl alcohol employed must be quite

The former product as asponshed by bosings with datas hydrochions and, which correct is time scetce near and methyl-indines hydrochiond. Another method of separating sunline from its mono- and di-methyl-derivatives is referred to in the feorhoods on page 76 Methyl-andiane can be re-founded by treating its nutroe-derivatives with its and hydrochions can't design the sunline state of the second state of the second second

³ The arbite must be fee from telumban and majorities resultable in Apricechiors each, and the methyl alcohed embeyed must be quite free from ethyl alcohed and accesses, the latter of which not only reduces the public but gives a product unvestibility for the preparation either of methyl valot or maintaints green, owing to the formation of a bias of the formation GLEGLE, NGCLA, 30 april or anime are used, of which it is on saturated.

free from ethyl aleohol and acetone, the latter of which not only reduces the yield, but gives a product unsuitable for the preparation either of methyl-violet or malachite-green, owing to the formation of a base of the formula —CH₂(C₂H₁ N(CH₂)_{b)s}

Dimethylamine is a colomiess city liquid, solutiving at 0°5 and boiling at 192°. It has a sharp base codour, and forms uncrystallisable saits. It untes with methyl icolde, with energy at the ordinary temperature, to form the todde of timethylphenylamiconium, which breaks up again into its constituents on distillation, but by reactions with argentic exide yields trimethyl-phenylamiconium in ydroxide, McgPhNOH, a crystalline, very deliousessenk conceive, and very bitter bases.

With bleaching-powder solution, dimethylaniline merely gives a pale yellow coloration, a teaction by which any contamination by anthre or mono-methylamiline can be detected, as these bases give a violet colour with the same reagent (page 45) Mild oxidising agents, such as chloranile, carbon oxychloride, and cupie chloride, convert the methylaniline into methyl violet (Part I page 234) With acid chlorides and aldehydes, it yields complex compounds Thus with benzaldehyde it gives tetramethyl-paradiamido-triphenylmethane, and the corresponding hydroxide or carbinol, Call [N(CHe)ele OH, obtained from this by exidation, is the base of malachite or benzaldehyde green (Part I page 241) By reaction with diazobenzens chloude, dimethylaniline is converted into dimethyl-amidoazobenzene, C.H. N. C.H. N(CH2), or butter yellow, while with diszobenzene-sulphonie acid it yields helianthin or methyl-orange (Part I page 188)

Paranth oco-dimethylamiline, CaH4(NO) N(CH3), is produced by the action of initite of sodium or initite of anyl on dimethylaniline. It is manufactured on a large scale for the production

with hydrochloric and and 75 parts of methyl alcohol The excess of methyl alcohol, and companisative jamil quantity of hydrochloric scal, tool to produce a purer oil With more hydrochloric scal, the restore takes place at a love interpretative, but there is a danger of forming following. The mixture is heard of first to a temperature of 270°, at a pressure not crossday 27 atmospheres When the teachers is complete, in about 15 hours, the pressure decreases without the temperature being reduced (Schoop, Ohen. Zet, xz. 125; 120° are Sec. Chem. 140°, r. 430)

¹ Ten put of dimetily-lamline are dissolved in 50 of strong hydrochlore and and 200 of water, and to the cold solution is gradually added a solution of 57 parts of solution mixture in 200 of water, when the hydrochlorate of the mixtuse-compound is oblistical as a body cayatilanging nyellow needles, from which the free base is obtained by treatment with potessuum carbinate and solution in other.

of methylene-blue, indophenol, and toluylene-red (Part I. pages 258, 285) It crystallises in large green plates or tables, soluble in other By exidation with potassium permanganate or ferrievanide, it is converted into paranitro-dimethylaniline, C₆H₄(NO₂) N(CH₅)₂, which forms long, sulphur-yellow needles, melting at 162°-163° When boiled with caustic alkalı, nitrosodimethylandine is completely split up into dimethylamine, H N(CHe), (which may, by this icaction, readily be obtained pure), and nitiosophenol or quinonoxime, C.H.O(NOH) (Part I page 157)

Commercial Dimethylandine usually contains more or less aniline and monomothyl-aniline By the entrance of methyl into the benzene-nucleus, more or less dimethyltoluidine, CaHa(CHa), N(CHa), and higher homologues are usually present in addition Hence the dimethylaniline of commerce usually boils between 198° and 205° The smaller the

range in the boiling-point the better the sample

The presence of aniline and monomethyl-aniline is indicated by the rise of temperature produced on treating 5 cc of the dry oil with an equal measure of acctic anhydride. This is stated to be 0° 815 C for each unit per cent of monomethylamine present. For small percentages this appears to be fairly correct, but with a product actually containing 30 per cent, an excess of over 7 per cent is said to be indicated. A serious objection to the method is that it wholly fails in presence of aniline. But the piesence of andme can be recognised by mixing a few drops of the oil with a few drops of ether, and adding one drop of strong sulphuric acid, when, it aniline be present, its sulphate will separate as a white precimtate

A more plausible method is that of Nolting and Boasson (Ber. x. 795), based on the different behaviour of the bases with nitrous acid, but the results yielded in piactice have been found

When annine hydrochloude is treated in cold solution with sodium nitrate. it yields diazobenzene chloride, while dimethylambne is converted into the hydrochloride of its nitroso-derivative (page 75) Both these bodies are freely soluble in water, while monomethyl-aniline is converted by the same treatment into the non-basic methylaniline-nifrosamine, which can be extracted by agutating the liquid with other. If this reaction occurred in its simplicity, the monomethyl-aniline could be estimated from the weight of the nitrosamine left on evaporating the ethereal solution. But when this is distilled in a current of steam, in which the nitrosamine is volatile, a considerable quantity of nitrophenyl-methylnitrosamine, C. H. (NO2) N(NO)(CH3), remains as a residue. This body is clearly produced by the oxidation of the nitrosamine, and direct experiment shows that pure monomethyl-andme, on treatment with excess of introus acid, is converted

unreliable by Reverdin and de la Harne. These channels recommend (Ohem. Zeit, xm. 387, 407; Jour. See Chem. Ind, vni 84), for the estimation of the aniline and methyl-aniline conjointly, activities too of the bases, and estimation of the excess of eactic anhydride by tiration with alkali and for the estimation of the aniline, dissotising and treating the product with beta-naphthol dissultoniae and

At ordinary temperatures acetic anhydride has no action on dimethylanihne, but on prolonged heating tetramethyldiamido-phenylmethaneis formed in considerable quantity. of the respent he in excess. Monomethyl-apiline is converted into methyl-acetanilide, C.H. N(CH.)(C.H.O), and aniline in the cold yields acetanilide, CaHs NHC2H2O, but on heating more or less discetanilide, CaH, N(CoH,O), is produced To avoid the formation of these secondary products the following method of working is recommended -From 1 to 2 grammes weight of the sample is mixed as rapidly as possible with an accurately known quantity (about twice its weight) of acetic anhydride, in a small flask fitted with a reflux condenser. After standing for half an hour at the ordinary temperature, 50 cc of water should be added, and the flask heated on the water-bath for fifty minutes to effect the conversion of the excess of acetic anhydride into acetic acid. The liquid is then cooled, diluted to a known volume, and an aliquot part titrated with standard caustic alkali. using phenolphthalein as an indicator. By this means the excess of acetic anhydride, CaHaOa, is ascentained, and the difference between the amount so found and that employed is the weight which has reacted with the ansline and methyl-ansline contained in the sample. 51 parts of acetic anhydride consumed in the reaction correspond to 107 of base in terms of methyl-analine, and the persentege of base thus found (a) is calculated and recorded

The ansime itself is determined as follows —From 7 to 8 grammes of the sample is dissolved in hydrochloric acid (28 to 30 a.c.), and diluted with water to 100 cc 10 cc of this solution

into it, to the exclusion of the simple unto same. As the molecular weights of the two bodies are materially different (181 186), the undefinite character of the reaction procents the accurate determination of the momently learning (Rover din and de la Harpe, Chem. Zeit, xiii, 387, 407; Jour. See Chem. Zeit, xiii 381)

¹ H. Giiand (Bull Soc Cham., 1889, in. 142) modifies this process by employing the acetic athydude dissolved in ten times its volume of dimethylamline. 10 cc of this solution is added to I gramme of the sample. After standing for one hour in a corbod flask, water is added, and the liquid (boiled for some time and) timited with standard benyive water or phenolythislein.

is further diluted with water and cooled by ice. The solution is then diazotised by adding a solution of solution into its in quantity sufficient to react with the whole of the sample if it consisted of aniline solely. A solution of the solution side between the cooled in the solution is sufficient to the solution side of the solution side of the solution side of the solution side of the solution is sufficient to the solution side of the solution is solution. As solution is sufficient to the solution is solution in the solution in the solution is sufficient to the solution is solution in the solution in the solution is sufficient to the solution in the solution is sufficient to the solution is solution in the solution in the solution is solution.

A measured quantity of this solution is then tessied with excess of solution actionate, and to it the second solution of the classification of the discourage of the second solution of the discourage of the second solution of the discourage of the second lall a prespirate coaces to form, when the liquid is filtered, and portions of the filtrate are tested with sail R and the discourage solution respectively, to ascertize which of these two is present excess. Another experiment is then made with suitably varied volumes, until after a few trains exact prespiration of the color volumes, until after a few trains exact prespiration of the color of disco-solution. The sections which occur are as follow.

$$\mathrm{C_6H_6\;NH_9HCl+HNO_2=C_6H_5N\;\;N\;Cl+2H_9O}$$
 , and

 $\mathbb{C}_u\mathbb{H}_p\mathbb{N}_2$ (21+C_nH_a(OH)(SO₂No)_a=HCl+C₂H_aN₂ C_mH_a(OH)(SO₂No)_b From those formule, and the volumes of the two solutions required for exact reaction, the weight of aniline present can be calculated. I gramme of sail E will react with 0 2672 gramme of sail is will react with 0 2672 gramme of sail is the found (b) is multiplied by $\mathbb{I}_{(a-\frac{m}{2})}$, which gives its equivalent in methyl-aniline, and this (c) subtracted from the sum of aniline and methyl-aniline in terms of methyl-aniline found by the acceptionation process (a) gives the percentage of real methyl-aniline (d) present. The dimethyl-aniline is determined by difference.

In the case of a sample of known composition, Reverdin and de la Harpe obtained the following satisfactory results by the foregoing process.—

	Procent.		Found.	
Amline,	10 42 1	oer cent.	10 30 p	er cent.
Monomethylaniline	, 1097		11 16	**
Dimethylandine (b)	y			
difference),	78 61	11	78.54	,,
· · ·				
	100.00		100 00	

The presence of monomethylandine is more objectionable in diamethylandine intended for the manufacture of green than in that to be used for violet. Schoop (Chem. Zeit., xi. 254) states that

the proportion seldom exceeds 2 per cent, and that the best qualities of dimethylandine are nonly or quite free from it. When 79 present, monomethylamine can be removed by shaking the oil with a small quantity of dilute sulphuric acid, or by boiling with acetic Diethylaniline C.H. N(C.H.)2

This base is best prepared by heating one molocule of audine hydrobromide with 10 per cent, in excess of one molecule of ethyl alcohol to 1±5° for 8 or 10 hours Nearly the theoretical yield is obtained The base boils at 213° 5 Diethyl-orthotoluidine and diethyl-paratoluidine may be obtained by oxactly similar means.

DIPHENYLAMINE PRENYLANILINE CoH, NH CoH, This base is obtained by heating annine with the hydrochloride or other salt of anilms 1 Diphenylamine crystallises in small white plates, having an agreeable flowery odour and burning tasto. It moits at 54°, and boils at 302° C (Graebe) It is almost insoluble in water, but readily in alcohol, ethic, benzeno, and aniline. Diphenylamine has very feeble basic properties. The hydrochlorule is a white crystalline powder, which turns blue in the air, and is decomposed by water The most characteristic reaction of diphenylamine is the deep blue colour produced by adding a trace of nitric acid to its solution in strong sulphuric acid. The reaction, which is very delicate, is employed as a test for nitric acid

Commercial diphenylamine should be pale yellow, melt not much below 54°, be free from unpleasant odour and oily matters, and give no violet coloration with bleaching powder for making diphenylamine blue, aurantia, and orange IV

Methyl-dynenylamone, CoH5 N(CH5)CoH5, 2 boils at 282°, and gives various colour-reactions with oxidising agents In dilute sulphurne acid it dissolves to form a liquid of the colour of solution of potassium permanganate.

I Six parts of aniline and 7 of aniline hydrochloride are heated to 250° under a pressure of 4 or 5 atmospheres for 24 hours
The ammonia formed is allowed to escape at intervals to prevent reconversion of the diphonylamine into amline. The product is treated with warm hydrochloric acid and a large quantity of water, which dissolves any anohanged aniline hydrochloride, and decomposes the hydrochloride of diphenylamine, which latter base separates

2 Made on a large scale by heating a mixture of 100 parts of diphenylanine, 68 of hydrochloric and (sp. gr. 117), and 2 parts of methyl alcohol for 10 hours, to 200° 250° at a pressure of 15 atmospheres. The product is treated with caustic sods, and the separated base distilled and shaken with twice its measure of strong hydrochlone acid The hydrochlonde of diphenylamine separates in the solid form, while that of the methyl-derivative forms a liquid, which is decomposed by adding a large quantity of water.

Warm nitric acid converts diphenylamine and its methyldorivative into $C_0H_2(NO_2)_k$ MH $C_0H_2(NO_2)_k$, hexanitro-diphenylamine, the ammonium salt of which constitutes the colouring matter known as assantia (Part I page 156)

Para-amide-diphenylamins issults from the reduction of phenylamide-azobenzene, nitro-phenylamine, or troposlin OO (Part I.

pages 181, 189, 190, 213)

TRIPHENYLAMINE DIPHENYLANILINE (C.H.) N

This body is formed by the action of brombenzene on dipolarsium anhine. It is a neutral body, melting at 127°, and crystallassiguing from other in monoclinic pyramids. It forms no isonitrile, piciate, nor acetyl-compound, but yields nodule of triphenyl-methyl-ammonium on treatment with methyl iodide. It solution in glacial acetic acid is coloured green on adding a little mirrs each, but with subburne and it gives a violet coloration changing to blue

Amidophenols,

By the reduction of the introphonois, corresponding amidocompounds are obtained. These bodies may also be prepaied by heating either of the three isomoric amido-hydroxybenzoic acids, C,H_(NH_1)OH COOU, with causite bayta

In the amulophenois the and character of the phenois m noutralned by the presence of the amulo-groups, so that they only yield salls with acids, but as phenois they are still capable of yielding alky-i-denvatives (eg., anisidine), while the hydrogen of their amulo-groups may be replaced for acety, &c, se in pla na cetin

The amulophenois form colourless crystaline scales or pites, which are very readily oxidable on exposure to an; with blackening and formation of resiones products, especially in impure. On the other hand, their hydrochlories are relatively stable, and often capable of sublimation. The solution of puramitophenoil hydrochlories in coloured first violet and then green by solution of blacking powder, qui in one och lorin inde, C₂H₁O(NCI), being formed, while with chromic and mixture, and other oxidising agents, it yields qui no ne, C₂H₂O₂. Treatment with subpluretted hydrogen and forme chloride converts to into compounds of the medigleme-blow group (Part I page 285)

The formyl- and acetyl-derivatives of the amidophenols are converted with great facility into anhydro-bases. Thus of henylamidophenol, a best liquid boiling at 200° to 201°, is obtained by boiling ortho-amidophenol with aceto anhydride.

When this body is heated with dilute acids, the reverse action occurs a cetyl-orthogomidon hen of being formed.

The methyl esters of the samlophenois (anisdines or samdo-anisols), and the corresponding ethyl esters (phenethidnes or antiolophenoids), are bases resembling aniline, and aremployed for producing certain soc-drys (eg., anisol vol., phenoids red, Pati I page 192) The acetyl-derivatives of the sesters are used in medicine under the names of instacctis and interporting less below?

The following table shows the characters of the isomeric amidophenois and their derivatives —

				_			-			
					ORYF	0 1 2	YES	4-1 8	PARA	1 4
					Melteng-	Boiling Point	Melting-	Boshing	Metteng. Point	Botteng-
Amidophenol (page 80), . C ₅ H ₄ { OH NH ₂					170	sub Itmes			184	•
Acetyl desivative (page 30), C6H4 (NH(COCH2)		•	•	•	201				179	
Mothyl ester (Amsidine), $C_6H_4\begin{cases} O(CH_3) \\ NH_4 \end{cases}$	٠	•	•	•		208		251	56	246
Ethyl ester (Phenethidme), $C_6H_4\begin{cases} O(C_2H_6) \\ NH_3 \end{cases}$		•	•	•		220	0	190-205 (at 100 mm)		253
Methacetin (page 85), $C_6H_4\begin{cases} O(CH_6) \\ NH(COCH_6) \end{cases}$	٠	•	٠	•	84	204			127	
Phenacetta (page SI), OgH (O(O)Hg) NH(OOCHg)	•	٠	•		70		97	-	135	
Amidophenacetin. Phenoc C ₀ H ₄ { O(C ₂ H ₅) NH(CO CH ₂ NH	oll (a)	(page	85),						100 5	

PHINAGRAINS ACES-PITENSTRIDINGS

The bodies of this formula have recently acquired some reputation as antipyretics and analgesics.

The phemicotins are prepared by ethylating the corresponding mono-mitrophenols, thus obtaining the isomers of the formula $C_0H_c(NO_2)$ OC_0H_6 On treatment with zinc or iron and hydrovol. III. PART II.

chloric and, these are reduced to the corresponding phenethiclines, $C_0H_4(\mathrm{MH}_2)$ OC_0H_3 , which are purified and non-tylised by heating with glarnal acetic and for some hours, the products being recrytallised from water

Of the three isomeric phenacetins, the meta-compound is unimportant. It forms tasteless and odourless scales, melting at 96°

Para-accelphenethaline is the official variety in the German and britch Plan manopiessas (1890). It forms withis, odouriess, insteless, gisterning scaly crystals. It requires 1400 pairs of cold, or 70 parts of boling, water for solution, and is soluble to a notable extent in chloroform. Its solution in 16 parts of alcohol is precuptated by the smallest addition of water. The crystals mids 1435*

Orth-acetphenethaline forms brillant white, very light spangles, without tasts or odour, and melting at 70°C I as very slightly soluble in cold, but more reachly in hot, water, separating again on cooling. It dissolves in about three pairs of rectified spirit, and abundantly in oblooform

Besides the differences in their melting-points and solubilities para- and ortho-phenacetia nar distinguished by their behaviour when boiled for several hours with didute sulphune seed (sp gt. 126). When thus treated, the para-compound yudds acche and and spainigly soluble sulphate of phenethidina. Orthophenacetin, on the other hand, is not decomposed by the same treatment, requiring the action of acud of 1.676 specific gravity for two home at 90° to effect its supomification. If in either case the acid liquid be disablesed, and then treated with an ammoniacal solution of naphthol-disniphonic acid, a fine red-yellow colour will be obtained if paraphenacien was employed, while with the ortho-compound a cherry-red coloration is produced. In either case the coloning matter may be precipitated by hime.

This formation of an exo-colouring matter may be employed to detect the phenacetims in truns and other organic liquids. The urns is evaporated to dryness, and the residue tested with hot alcohol. The solution is discord, evaporated, and the residue boiled for two hours with dilute sulphunc and (sp gr. 126) under a refux condensor. The resultant solution is cooled to δ' or δ' C, tested with a 1 per cent, solution of solution interior for five minutes, and then poured into a solution of naphid-disulphone and in excess of ammons, taking care that the mixture remains

¹ S Lutkke detects orthophemeath by boling 15 grammes of the sample with 25 grammes of dilute hydrochions and, when ortho-phemethican hydrochloude as formed, from which the fice base may be separated by caustic sola, and its boling-point (given by Lutkke as 247 5) determined. The hydrochloude gives a blood-red coloration with ferric chloude. alkaline If either modification of phenacetin be present in the urine a characteristic coloration will be produced, from the intensity of which the amount of phenacetin may be estimated

For medicinal use, phonacetin is said to present considerable deviatations over antitybrine, and especially over antitybrine (acetanitide), for while the lattet body is decomposed in the system with formation of an line, which has musked toxic properties, phenacetin yields phenacetin the control of an line, the control of the control

According to Revier (Pharm Zest, 1891, page 185) phenacetin is hable to contain unconveited prace-phenelulariae, which appears to be posenous in very small closes, if taken for some time, producing nephritis and albuminum. To detect the impurity, leuter meits 2½ grammes of chloral hydrate at 100°, and adds 0 5 gramme of the sample On agitation the phasicieta dissolved, and, if pure, the solution will regular cloudless when heated on the water-bath for five munites, though after longer heating it will assume a ruse tint. In presence of prare-phenchium, an intense coloration, ranging from red-violet to blue-violet, is produced in two or three munites at most.

S. Luttke detects diamidophenols or diamidophenatols in phonacetan by grading 0.5 gramme of bleaching powder to a fine paste with hydrochloric acid, and adding about 0.03 of the sample, when a red colour will be produced.

The lower price of aectanitals, and its close physical resemblance to phenoscin, have suggested the possibility of the partial or only please abstitution of the former body for the latter, and a figural instance of such a practice is actually on record (Pham Jour. 18), xxi 377). The presence of 5 per cent of acetanithe lower the melting-point of the sample to $127^{n}-128^{n}$

H. Schwartz (Phanm Jour., [3], xum 1085) recommends that I gramme of the suspected sample should be heated with 2 c.c. of causts coda solution, a fragment of chloral hydrate or a few drops of chloroform added, and the mxture again gently heated With phenacetin the odour is aromatic and not dissgreeable, but in presence of acetamilide, the penetiating and repulsive smell of phenyl-arbamin, C.H. NG, is produced. On boiling the asmple with caustic soda solution, only deep of antime separate if acctamilide be present in considerable quantity. If the cooled liquid, together with the separated globules, be shaken with either, and the other separated and evaporated, the resulce when tissolved in water and treated with a drop of carboic soul, and a clear solution of bleaching powder saided, gives a blue-green coloration changed to omore—d by hydrochlore said, and restored by ammonia (Soe also Jour. Soc Chem. Ind. vin. 72).

For the detection of so-taminds in phenacetin, M J S ch r o der in recommends that 0 8 gramme of the sample should be bouled with 8 cc of water, and the input filtered whose old from the recystallised phenacetin. The filtrate is bouled with a little potassium nitrate and didute nitro eard, a solution of mercurous minate containing a little nitrous and added, and the whole sigms houled. A red colour will be obtained if the proportion of acstanlishe in the sample accords 2 per cent

If I gramme of a muxture of equal parts of phenacent with acetanihib es baikaen with 200 cc of water, the whole of the acetanihide goes into solution together with 0130 gramme of phenacetin, whils the remainder of the phenacetin remains miscluble. If this be separated, its weight, when corrected by an addition of 0130, will represent the phenacetin present in 1 gramme of the sample (Pham Jour., [3], xu. 377)

Phensestin has been made official in the German Pharmacognia (1890), the maximum does being I grainine. It is stated to melt at 135°, and dissolve in 1400 perts of cold, 70 of boiling, water, and 16 of spirit to form neutial solutions. It is dissurguished from exalgen and antiebrin by boiling 01 graume for a munite with 1 ec of hydrochloric said, adding 10 ec of water, filtering, and adding to the filitate 3 drops of a 3 per cent, solution of chronic said, when a ruby-red colour will be gradually developed. (See Pharm Jour. [3] Xxx 1978) Strong sulphura said should dissolve phenacetin without becoming coloured, while a saturated solution, if free from phenol and acetanilde, will not become turbid on adding bromine-water. The description of phenacetin in the British Pharmacognesia additions (1890) closely corresponds with the above. The does is from 5 to 10 grains.

Mchip-thesacetra, C,H_(O C,H_) N(CH_)(C,H_C). This body is prepaid by testing para-pleasectin un xylene solution with sodium, and causing the resultant sodium-derivative to nect with methyl iodde (PAram Jons, [3], xxi 81). The new product distils at about 300°C as an oil which crystallines on standing. It may be purified by recrystallisation from alcohol or ether, when it forms colouriess crystals, moderately soluble in water, and having marked narcotic as well as antipyretic characters

Amido-pas aphenocates, C.H.(O C.H.) NH(O ČH ML) The Apidrochloride of this base is readily soluble in water and alcohol, and has been introduced, under the name of "phenocollum high ochlorioum," as an antipyretic and antirheumatic Prolonged boiling with alkales splits it into pare-phenethichine and glycocine.

Formyl-paraphenethidene, $C_6H_4(O\ C_2H_5)$ NH(CO H), though having a constitution similar to acet-phenethidine, appears to have no antipyretic properties, but has been suggested as an antidote in cases of poisoning by strychinne

Methacetin is the commercial name of para-acet-anisidine, CoH4(O CH2) NH CoH2O. It is, consequently, the lower homologue of phenacetin (page 81). It forms a crystalline powder or small lustrous scales or plates, odourless, but of a faintly bitter taste It melts at 127° C, and at a higher temperature boils and distils unchanged. It dissolves in 526 parts of cold, or 12 of builing, water, and is easily soluble in alcohol, acctone, chloroform, and dilute acid and alkaline liquids. It is less soluble in benzene, and only with difficulty in ether, carbon disulphide, petroleum spirit, and oil of turpentine, but dissolves freely, on warming, in glycerin and fixed oils. In its general reactions and physiological effects, metacetin closely resembles phenacetin, though according to some authorities it has a less powerful, and according to others a more powerful, action. Its efficacy in cases of neuraliza and rheumatism is said to greatly exceed phenacetin, from which it may be distinguished by its physical characters, or by heating it with a quantity of water insufficient for its solution. When thus treated, methacetin melts and solidifies again on cooling, whereas phenacetin undergoes no apparent change 1 cc, of hydrochloric and dissolves 01 gramme of methacetin very easily, whereas the same quantity of phenacetin is mainly undissolved

DIAMIDOPHENOIS C.H. (OH)(NH.)

These bodies are weak bases, forming salts which crystallise well and give aqueous solutions which turn brown in the air, and are coloured an intense violet or dark red by potassium bichromate, ferric chloride, or blenching powder

TRIAMIDOPHENOL CaHa(OH)(NHa)

This body is an unstable base resulting from the complete reduction of picric acid, $C_0H_2(OH)(NO_2)_n$ med solutions. If alkaline reducing agents be employed, the action does not proceed beyond the formation of $\dim \operatorname{Iniro-amid} \operatorname{do-phenoic}$ picra mic acid, $O_2H_2(OH)(NH_2)(NO_2)$, see Part I, page 143)

A dilute solution of triamidophenol is coloured deep blue by

Phenylene-diamines. Diamidobenzenes.

Three modifications of phenylene-diamine or diamido-benzene, $C_0H_4(\mathrm{NH}_2)_2$, are known, differing from each other in properties according to the positions of the amido-groups, thus —

	Ortho Compound 1 2	Meta Compound 1 8	Para Compound
Appearance, . Melting point, Boiling point,	Tablets or plates 102°-105° 252°	Crystalline mass 68° 287°	Tablets or small plates 140° 207°
Characters of hydro chloride,	Groups of radiating needles, readily soluble	Concontilically ar- ranged crystals	Readily soluble tablets, very spatingly soluble in hydrochloric acid
Reaction in neutral solu- tion with sodium nitrite,	Separation of amido azo phonylene as a colouriess ofly liquid	Yellow or brown coloration, on precipitate of triamidoaso benzeno	No rouetton

Ortho-Phenylens-diamne is distinguished from its isomendes by its reaction with sodium nitrite, and by the separation of unby-red needles on aiding ferric chlonds to the solution of its hydrochloude. On treating an alcohole solution of the base with a drop of phenanthraquinon dissolved in glacula sceles each, and boining for a short time, a bright yellow prompitate of diphenylens-quinoxalinc, Cophyn, is formed. It consists of ampil needles which are coloured a deep red by strong hydrochloric and, and its production affords the most delicate reaction for ortho-phenylenschamme. Its somewise one of the reaction, but its homologue, ortho-toluylenediatine, therefore the reaction for other seminative.

MERI-PERNYLMEN-DIAMINE may be prepared by the reduction of meta-dustrobeause (Part I page 178, footnote). It often semans in a state of superfusion for some time, but is instantly solidified by adding a crystal of the solid substance. Metaphonylene-dimining is sparingly solidide in water, the solidion being alkaline in reaction. It is reachly solidide in other, and may be extinated by this solvent from alkaline aqueous liquids. It is a di-eard base, the hydrochloride bound G.H.(DHL, p.HCI. The reaction of metaphenylenetical with solidium nutries is characteristic and extremely delicate. It is due to the forestation of Bismarch's or phasulace brows (Part II page 180), and by means of it one part per million of introus acid can be detected in water

Metaphenylenedamme possesses maked possonous properties, its physiological action resembling that of the leucomannes and phomaines. Du bois and Vignon (Compt. Rend., vois 533) experimented on dogs, and found that a dose of 0.1 gramme per kilogramme of the animal produced salvation, vomiting, diarrhos, abundant excretion of urne at intervals, and death by come in twelve to fifteen hours. Besides these severe symptoms, all those of intense influenza were produced, such as acute coryza and sneezing, couchine, and extreme decreases.

Park-Thenyless-dualing occus in anilms tailings (page 67). It may be prepared by the reduction of paramitanetenilule. It is but slightly soluble in water, but readily in alcohol and other. When heated with dilute sulphune and and mangamese duzule is plus (q u in on e, G_pH₁O_p, which reaction distinguishes it from its isomerides. On passing sulphunetted hydrogen through a solution of the hydrochloride, and then adding forms chloride, throwne or Lauth's model is formed (Part I was 285).

Para-phenylenedmanuse possesses possonous properties similar to those of meta-phenylenedmanus, but death occurs more rapidly than with the latter base. It also exerts a special action on the eye, which is gradually forced out of its othic by the swelling of the conjunctiva or inter-orbital cellular tasses, while the lachrymal glands are blackened by a deposit of pigment (compare "Tolivylenedwanuse")

Dimethyl-paraphenylenediamne, H.N.C.H., N(CH₂), may be obtained by the reduction of introcodumethyl-anilme or delitabilities (Pat I pages 188, 211) A neutral solution of the hydrochloride is coloured a beautiful purple by force holonde, and on treating it with a hydrochloric and solution of sulphuretted hydrogen, and then adding ferric chloride till the smell of sulphuretted hydrogen, has disappeared, a fine blue coloration is obtained, due to the formation of methylene blue (Part I page 285). This resident is the most delibetic test for

TOUTLENS-DIAMNESS DIAMNOTOLDENSS. C.J.H.(CH.N.N.L.).
These bases closely resemble the phonylene-diamnes. The orthopara-modification (CH., NH., NH., = 1: 2 4) is obtained by the
reduction of ordunary diamteriolismes. It melts at 88°, is used for
the production of tolugiane ved and tolugiane orange. The 1: 3: 4
(meta-para) modification is obtained by mitrofying seel-parasionized
seporitying, and reducing 1 3 is no vs ky (Jone. See Oftem Ind.,

¹ This modification appears to be identical with the paratolnylenediamine isolated by Hell and Schoop from aniline tailings (Berickle, Mr. 723).

1x 383) gives the following table of reactions of neutral or slightly acid solutions of the two isomeric toluylene-diam nes -

Reagent	α-Toluylene-diamine CHg NH ₂ : NH ₂ =1 2 4	β -Toluylone-diamine CH_0 NH_2 $NH_2=1$ 3 4		
Ferrio obloride	No change at first, after standing for a long time an orange coloration	Wine-red coloration		
Potassium bichromate	Yellowish-brown colors-	Reddish-brown precipi-		
Potasshua ferricyanide	Olivo-green crystalline plates	Dark red coloration.		
Bromine water	Yellowish white precipi-	Brown flocks and magenta red solution		
Platinic chloride	Yellowish-brown colora tion	Reddish-brown precipi-		
Aurie chloride	Brown proofpitate	Red solution with blue reflex and motalite mit- ror in the cold		
Potazsium nitrite	In very ditute solutions a golden-brown coloration, in concentrated a brown precipitate	No coloustion, but a sal mon-coloured precept- tate		
Solution of bleaching powder	Reddish - brown colora- tion and then a light brownish yellow precipi- tate	Dark-red coloration, then an clive green precipi- tate		

The foregoing reactions are available, even in presence of other substances, for the detection and identification of the toluylenediamines, which often result from the reduction of azo-dyes.

The toluylene-diamines are powerful poisons (compare "Metaphenylenediamine," page 87)1

Benzidine. Dipara-amido-diphenyl.

$$C_{12}H_{12}N_2 = NH_2C_6H_4C_6H_4NH_2(1, 4: 1, 4)$$

This body is obtained by the reduction of diparantico-diphenyl, NO₂ C₀H₂C₀H₂C₀H₃CO₉ by nascent hydrogen (an and hydrochloric aud) A readier method of preparation is the following —An alcoholes solution of 10 parts of asobenzene, C₂H₃N N C₃H₄.

¹ Engel and Kiener (Greigt Rend, vr. 465, Jour Clam Sec, hr 6) in this symptomic to vary considerably occording to the time required to the produce doubt, which ranges from a few bours in earls cases to servand weeks in chorne cases. When death reasses in a few days, there is always interior doctrone cases. When death reasses in a few days, there is always interior doffen hemoglobinums, and the urmo is loaded with this and yallow and brown regiment granules, which sometimes contain rom a Time foreignous product and account of the contain row is considered in the product of the contain row of the region in the procephase from the solubles, and not from the red corruption in the procephase from the solubles, and not from the red corruption.

is treated with a solution of 31 parts of tin in concentrated hydrochloric acid, and the hauld warmed for some time Hydrazobenzene, CaH, NH NH CaH, as formed, which by intramolecular change is converted into hence dine (dibulno-Some of the moment ortho-para-diamidediphenvl is simultaneously formed, and a portion of the azobenzene is reduced to aniline, CaHaNHo. The alcohol is distilled off the residue dissolved in water, and sulphime and added The nearly insoluble benziding sulphate is precipitated. while the sulphates of the isomeric base and of aniline remain in solution. The mecimitate is washed with dilute hydrochloric acad (to remove tin salts) and treated with ammonia, the bberated benzidine being crystallised from dilute alcohol. Benzidine is also produced by treating azobenzene with subbur diaxide Benzidine is manufactured on a large scale by heating nitrobenzene with caustic soda, a little alcohol, and the proportion of zinc-dust themetically sufficient to reduce it to hydrozobenzene The product is washed with cold dilute hydrochloric and to remove exide of zenc. On subsequently heating it with dilute hydrochlonic acid, it is converted into benzidine dihydrochlonide.

Benzadine forms large pearly plates, which are colourless when pure, but rapidly turn red on exposure to the air. It melts at 122°, and boils with pertial decomposition above 360° Benzadine is very sparingly soluble in cold but readily in bothing water, and

as easily soluble in alcohol and other

Bennádne vsa well-defined di-sad base, forming crystallashle sellar. The sulphate as very sparnigly soluble in water, even when boiling On adding potassuum behromate to a concentrated solution of beandine hydroboliorde, a deep blue crystallan preepitate. On taming C_pH_c(NH_p)_cCO_d, is immediately formed. The same necessitate is formed on warming, even in very dulue solutions.

When chlorme-water is added in small quantity of a solution of benatine hydrochlorids, the liquid assumes a fine blue colour, which on further addition of chlorme-water changes to green; and ultimately, when the chlorme is in excess, a focusion of precipitate is formed, apparently containing C₁₂E₁C₁₂N₂O₃ soluble in alcohol and ether, and forming a colourless compound on reduction Bromine-water and a solution of blueaching bowdes at similarly, but in presence of a large quantity of free hydrochloric and bromine forms terts about the native liquid production and bromine forms to tert about the native late, making at 285° With introve and, solutions of benzeline salts react to form terts are or in pour and such in the control of the colour production of

which congo-red is the type (Part I page 206), and which are remarkable for dveing cotton without a mordant

ORTHOTOLIDIEN NH₂ C₆H₂(CH₂)(CH₃)C₆H₃NH₃ This base is homologous with benadue, and is prepared from ortho-nitro-tolusne by the same process by which bonadune is spepared from introbenzene. It melts at 128°, and presents a close recemblance to benadume. The text is a -d ye s prepared from the rales readily altered by audit than are the similar dyes prepared from benadine.

NAPHTHYLAMINES AND THEIR ALLIES.

When naphihalane, $C_{ij}H_{ij}$ is treated contously with intro and, nitronaphibalane, $C_{ij}H_{ij}(NO_j)$, is formed, and this by testiment with reducing speats is converted into an ido-naphihalane or naphihylamine, $C_{ij}H_{ij}(NH_{ij})$. These rescens are stretly unalogous to those by which aniline is prepared from benzene, and the product is known as alpha-naphihylamine by other resolutions the isomere betwin aphihylamine may be obtained. These two bodies differ from each other na notable manner, as unducated in the following table—

	Alpha Naphthylamine.	Beta- Naphthylamine
Structural Formula, .	HO CH CH	HO CH OH OH
Melting-point,	50°	113
Boiling-point,	800°	294°
Odour,	Dragreeablo, persistent	None
Appourance, .	Plat neodles or pusms	Pearly plates
Reactions of hydrochlo- ride in solution — With ferits chloride, With introos acid in alcoholic or nestic acid solution, With sulphambra acid and solution mitrie, followed by hydro- chloric acid,	Blue precupitate Yellow colour, tuined crimson by hydro- chlone acid Red coloration	No reaction No reaction

a-Naphthylamine. CiaH, NH.

This base is obtained (as aheady stated) by the reduction of nitronaphthalene, or by heating a-naphthol with the double compound of chloride of calcium and ammonia.

- a-Naphthylamine has a most disgusting and persistent odour, resembling that of faces: It turns volet or brown in the air, but when purified by sublimation this change occurs very slowly, and only on exposure to an and hight It is slightly volstile with steam.
- a Naphthylamine is nearly insoluble in water, but very soluble in alcohol and ether. It forms a series of readily-crystallisable, easily-soluble salts. On adding ammonia to a solution of the sulphate, the free base is precipitated in white silky needles.
- On adding ferrse chlords to a solution of e-saphthylamme, or of one of its salts, an arms bits precipitate of n a ph it has me in ment with sulpharous and Other outlining agents (e.g., chomor each, bleaching powder) produce precipitates varying in colour from blue to violet or ord?
- On adding an alcoholic solution of introus acid to a solution of enaphthylamine in alcohol or glanial sectic acid, a yellow coloui is produced, which, on adding a little hydrochloric acid, changes to an intense violet or magenta colour, or, in presence of only traces of naphthylamine, to a reddsh colour

If to a cold solution of alpha-naphthylamine sulphanilic acid and solution nitrite be added, a red colour is produced on adding hydrochloric acid, owing to the formation of a mid on a pht hyl-

- a zo b e n z e n e-s ul p h o n 1 c a c 1 d, C₁₀H₆(NH₂) N₂ C₆H₄(SO₆H)
 a-Naphthylamine is used for the preparation of Magdala ed (Part
 I p 257), certain azo-tlyes, and naphthalene faucy-colours on cotton,
- Commercial a-naphthylamine ought to melt at 50°C, and be almost completely soluble in dilute hydrochloric acid. Naphthalene, the presence of which causes incomplete solubility, may be demined by distilling the sendulated solution in a current of steam, agitating the distillate with ether, separating the othereal layer, evaporating it at a low temperature, and weighting the residence
- 1 On a large coals, a-saphthylamna s prepared in a manner very similar to that amplied for the production of antine. Nutrosaphthalene is reduced by ron and hydrochlouse and at a temperature of about 50° When the reduction is complete, milk of time a solded, and the maphthylamna distribution, when it is obtained as a nearly colouries onl, which solidiles to cystalline cakes of grayant colour. It appears to be wholly free from p-maphthylamne, but contains an impunity which is probably 1 1'-n a ph th ylene-diamine, Culture (Laglen Mark) (O. Witt, Dang Sight, Jour, colour 255).

B Naphthylamine. C,H, NH.

This modification of amidonaphthalene is most readily obtained by heating β -naphthol under pressure with ammonia at 160°, or with the double compound of zinc chloride and ammonia at 200°-210°.

Becomes compound to the control of the manifest of the than the α -nodification. It volatheses in a current of steam, and is alightly soluble fluorescence, which, however, is not shown by β -naphthylamine salts β -Naphthylamine gives no coloration with oxidiang agents, nor with introls and hydrochrone acids in allocobes solution.

Commerciat β-naphthylamine ought to melt at 112° C, and be completely soluble in dilute hydrochloric acid

Tetrahydro-β-naphthylamine C₁₀H₁₁ NH₂

This base has been introduced into machinic under the name of "The run ine" it is a colouries, algibity viscous liquid, of perchiar cdour. It is a strong base, a drop soon becoming converted into a crystalline mass of the corboxide on exposure to air. The hydrochloride forms well-defined white crystals, incling at 237°, and readily soluble in water, alcohol, and amply alcohol.

The physiological effects of thermine embrace the two strongly-marked characteristics of mydrasis (accompanied by pain) and elevation of the temperature, which latter effect has been observed to the extent of 44° C

Naphthylamine-Sulphonic Acids.

When treated with dilute sulphuric acid, the naphthylamines dissolve easily with formation of sulphates, but by the action of concentrated sulphuric acid at a high temperature they are converted into sulphonic acids Thus when α-naphthylamine is heated with furning sulphuric acid, two isomeric sulphonic acids are formed, one of which is readily soluble in water, while the other is only sparingly soluble. The latter modification crystallises in small lustrous needles, and in aqueous solution exhibits a beautiful fluorescence. Similarly, β -naphthylamine yields on sulphonation several momente acids According to A G. Green (Bé) , xxii 721), at moderate temperatures (100° C), and with ordinary sulplume acid, the product is a mixture of a and y acids, having their sulphonic groups in the a-position, while at a higher temperature (160°-170°) B and S modifications are produced, having their sulphonic groups in the β -position. The ammonium salt of the β -acid is less soluble than the three isomeric salts, and by this means the β -acid can readily be isolated.

The α -naphthylamine-sulphonic acids may also be obtained by treating initronaphthalene, $C_{10}H_{2}NO_{2}$ with fuming sul-

phuric acid, and reducing the resultant intronaphthalene-sulphonic acid, $C_{10}H_6(NO_2)(SO_3H)$, with iron and hydrochloric acid. Two isomeric annide-sulphonic acids are obtained in this case also

The naphthylamine-sulphonic acids are also conveniently prepared by heating the corresponding naphthol-sulphonic acids (Part I pages 194, 207, 208) with ammonia under pressure

Naphthylamine-disulphonic acids may be obtained by reactions similate those described above. Two of these derivatives of \$\beta\$-naphthylamine are technically known as "Amidoacid \$R" and "Amido-caid \$\beta\$. The latter, or \(\triangle \), \(\triangle \

Naphthylene-Diamines. C10H6(NH2)2

These bases may be formed by heating the corresponding dihydroxynaphthalenes with ammonia, by the reduction of the dimitionaphthalenes, and in other ways

The following table exhibits their leading properties -

Position of the Amide Groups	a1, 48	a ₃ , a ₅	a1, a4	α ₁ , β ₂	
Mode of preparation,	nambithyle miteo nanh nitro-nanh- azo-com- I		From a ₁ – a ₈ . Dihydroxy- naphthalene.		
Form of crystals,	Leuves	Noodles.	· Needles,	Plates.	Needles
Melting point,	120° C	189° 5 C	66° 6 C	95° O	189° C.
Hydrochloude,	Plates	Necdles (?)		Plates	Plates
Sulphate,		Needles		Pintes	Needles
Reaction of the hy- drochloride with femic chloride,	Green colora- tion	Blue coloru- tion, then blue precipa- tate	Chestnut- hown pre- cipitate	Green, then yellow coloration, brown pre- cipitate	Blue colora- tion, then precipitate
Action of nitrous acid,	Sol tetraro- compound	Sol tetrazo- compound	Vermilion precipitate	7	Sol tetrazo- compound
Action of the arc dye- stuffs on ununci- danted cotton,	Do not dye	Dye the fibre	-		Dye the fibre

¹ For further information respecting th and the neighthalene derivatives generally, and Wynne (Jour and Proc Chem So cle by

. Wynne in Thorpe's Dutionary of Applied Chemistry, in 649 et se

Amidonaphthols, C10H0(OH)(NH2).

These bodies are unstable bases obtained by the action of reducing agents on the intro- or introso-naphthols, or on certain azo-dyes. The following table shows the leading differences of the principal members of the group .—

	a-Amido- a-naphthol	B-Amido- a-usphthoi	a-Amido- g-naphthol.
Relative position of the OH and NH ₂ groups	1 4	1 2	2 1(or 4)
Mode of turnfation	Reduction of 1 4 nitro s naphthol melting at 161°, or of Orange I (Part I page 284)	Reduction of 1.2 nitro - α-naphthol melting at 123°, or of nitroso α-naph- thol	Reduction of the nitro-β-naphthol, melting at 108°, of nitroso-β-naph- thol, or of Orange II (Part I page 184)
Characters of free base	Unstable,	Unatable	Colourless scales; elightly soluble in water, oxidesed in the sir Ethe- real solution ex- hibits violet fluo- rescence
Reaction on agretating alkalino solu- tion with all	Dhty green colora- tion, changing to yellow	Permanent graes- green cotour, and green scum sol- ubis in alcohol to pure green solution Orvioist naphthoquin- onlmide- CtoHe NH	Brown coloration
Reaction with bro- mine water	Yollowish-white needles precipi- tated, even in very dilute solutions	Yellowish or green preolpitate (the same with ferric chloride)	-
Characters of hydro chloride,	Long white needles or acteular plates with bleaching powder yields (29H2N2C), which separates from acutic acid solution in needles, melting at 85° and exploding at 189°	White lamine	White lustrous needles readily soluble in water, but only spar- ingly in dilute hydrochloric acid
Product of oxida- tion with chro- mic cold mixture	Theoretical yield of a-naphthsquinone,	β-naphthoquinons	β-nnphthoguinone

ANIDOMAPHTHOR-SUFFIGHTO AGDS. These bodies result from the reduction of ace-destratives of the respective disableness compounds of naphthol-sulphome ands. Thus, for metance, by treating the four known modifications of B-naphthol-monosulphome and with stannow chlords, O N. W. I. i obtained the following amidosulphome cands (Berichte, xx. 1468, 3489): —

- Amido-β-naphthol-β-sulphonic acid, from Schaffer's acid (Part I .page 194)
- Amido-β-naphthol-a-sulphonic acid, from Bayer's acid (Part I page 194)
- Amido-β-naphthol-δ-sulphonic acid, from Casella's acid (Part I page 208)
 - Amido-β-naphthol-γ-sulphonic acid, from Dahl's acid.

The first of these could has recently recovered a novel appheation as a photographic developer under the name of ellosopen (R M eld ols. Nour Sec Chem Ind., vin 885). It may be obtained by the sediction of the nead-ye known as "Crocem orange." "Brilliamt orange" or "Ponceau 4GB" (Part I page 184), obtuned by the reaction of Schaffie's 'B-naphthol-sulphone seni (Part I page 184) and inchember obligation in Imay be obtained from its interoclerivative by dissolving the ammonium or other salt of Schaffie's and in necodid water, together with an equivalent quantity of sodium ninite, and then gradually adding hydrochloric send to actif cachen, when the nitrose-and is at ones formed, and impairs an orange colour to the solution. The sand can be punified by conversion must be harmonic addinger to calcium salt (Gour Them Sec, XXXX 44), or the solution may be at once reduced to the similar-and by treatment with sun-duat or statamous chloude.

Two other am 1 do β -n ap h tholmonosulphone acids are obtainable by heating with caustic alkain, to 2007-280°, the two β -naphthylammo-insulphone acads respectively obtained by testing with the two isomerse β -naphthol-dasulphone acids R and Y (described in Emp Patent, 1878, No. 1715). They drifts from its amiltonaphthol-nulphone acids, referred to above, in yielding discompounds They can also be combined with various totrus-compounds, grung blacksh violeto blackship discontinuity and the second patents of the second p

	B	¥
Solution of neutral salts in water	Violet fluorescence	Blue
Reaction with ferric chlo- ride	Dark blue coloration, turn- ing to dun colour,	Durty claret-red coloration
Reaction with bleaching- powder solution	Light yellowish-brown coloration, which dis- appears rapidly on add- ing excess of the resgent	Dark reddish-brown colors tson, which disappears gradually on adding ex- cess of the reagent
Diago-compound	Reddish orange	Canary yellow.
Combination of the diszo- compound with "Realt" in an alkaline solution	Claret red.	Violet-black,

PYRIDINE BASES. C.H. N.

These bases, metamente with annihus and its homologues, are contained in coal-tar naphths, in shahe-oil, in pent-tax, in bobacco-smoke, and, together with aminoma and nethylamine and its homologues, in the product called "Dippel's oil," obtained by the distillation of bones and other amind mattics Pyuline itself has receaved several technological applications, and is of great interest theoretically in relation to the alkador to the distillation.

Pyridine may be regarded as benzene, in which one of the CH groups has been replaced by N¹ Thus.—

The homologous bases are derived from pyridine by the substitution of $CH_{\rm b}/CH_{\rm p}$ &c, for one or more of the hydrogen atoms, and consequently admit of isomeric modification according to the position of the substituted atoms in the chain

The following is a list of the bases of the pyridine series. The

¹ The relationship between various organic bodies (hypothetical and otherwise), of which the names commence with the root pyr is shown by the following formula (compare page 30). The hydrocathon pyrese has the constitution.
of a phenylene-naphthalene, and is not rolated closely to the bodies tabulated below.

Pisane has merely a hypothetical resistance, and the dihydride is known only through its diphenyl-derivative Pyrone and pyrazine, else, are only known by their derivatives Pyrozole, C,H,No, has been recently obtained by acting on hydrazine hydrate with emphelohydrin in presence of gno chloride.—

$$2N_gH_4 + C_gH_gClO = C_gH_4N_2 + HCl + H_gO + 2NH_g$$

Pyrazole is a basic substance orystallising in needles, melting at 70°, and boiling at 188°. It is readily soluble in water, alcohel, and other.

boiling-points and specific gravities are only approximate, as the isomeric modifications exhibit sensible differences in their physical properties

Formula	Base.	Booking-Point	Spensio at 0° 0	Gravity at 22' 0
C ⁶ H ⁶ M	Pyridine	115-116	9868	
C ₅ H ₇ N	Picoline	188-186	9813	*988
C ₇ H ₅ N	(e Methyl Pyridine) Lutidine	154	9448	ŀ
, CaHlon	(y-Ethyl-Pyridina) Collidine	179	921	
CoH13N	Parvoline	188	008	
C ₁₀ H ₁₅ N	Corridine	211	١.	974
O11H17N	Rubidine	280		1 017
C ₁₅ H ₁₉ N	Viridine.	251	١.	1 024

From the above table at as evident that the bothing-points rise as the number of carbon-atoms in the molecule increases. For the first four members of the sense the specific gravity diminishes, with increase in the molecular weight, but with the higher members the revease is recorded as being the case. The lower members are missible with water in all proportions, but collidine and its higher homologues are insolvible, or nearly on water.

If a drop or two of pyridine, or one of its homologues, be warmed in a test-tube with a similar quantity of methyl iodide, the product mixed with powdered caustic potash and moistened with water, and heat applied, a highly characteristic and peculiar odour is produced, owing to the formation of a pyridic dihydride. It resembles that of a mixture of mustard oil and isomitrile. The least trace of pyridine or its homologues can be detected in this way. A somewhat similar odour is obtained when a quinoline base is treated in the same manner, but the aniline bases and piperidine do not give the reaction The foregoing test, duo to A. W Hofmann, is modified by de Coninck as follows -1 c.c of the base is gradually mixed with 2 cc. of methyl rodide, the bround being cooled during the mixing. The crystalline product is dissolved in about 5 cc of alcohol, the liquid heated to boiling, and very concentrated caustic potash solution dropped in A blood-red colour is produced, and the liquid finally becomes dark brown if a pyridine base be present (Jour Chem Soc , 1 897) Piperidine, sparteine, cicutine, and the aniline bases give no similar reaction

The bases of the pyridine series are tertiary monamines, and

form with alkyl iodides compounds which are not decomposed by caustic potash, but yield caustic hydroxides by reaction with silver oxide (compare page 18).

The pyridine bases and their salts axer's a soporfic action on the higher animals. When inhaled, pyridine aste as a respiratory sodative. It has been successfully used as a beat simulant and as a topscal antespita mighthens. Pen ris held found pyridine to act as a general antiseptic, especially as regards magiciar. In the lower animals, pyridine and its homologues act as vulent poisons, and have been successfully employed in 0.2 per centrolium for destroying the scale-nears in sheep, the run-closse, and other injurious meets. The pyridine bases appear to be little, at all, infection to motion for these purposes, and thave also been employed in dissufficient powders.

For the preparation of the pyridine bases, bone-oil, or the fraction of coal-tar or shale-oil boiling between 80° and 250°, should be acutated with sulphuric acid diluted with twice its measure of water, the treatment being repeated to ensure the complete solution of the bases The acid liquid is separated and distilled (or boiled by a current of steam) till the vapours no longer redden a slip of fir-wood moistened with hydrochloric acid, showing that all the pyrrol has been driven off The liquid is then filtered through linen to separate tarry matters, an excess of caustic soda added, and the whole distilled with steam as long as bases continue to pass over, as indicated by the production of fumes by contact of the vapours with hydrochloric acid. The distillate is allowed to cool, and is then treated gradually with a large quantity of solid caustic potash or soda, till the pyridine bases separate as an oily layer on the surface of the alkaline lev 2 The upper stratum is separated, and, if it contains aniline, furning nitric acid is cautiously added and the mixture gradually heated to boiling, whereby the and ine is destroyed, while the pyridine bases remain intact 8 Water is then added, the precipitate filtered off, and the filtrate

¹ Their methiodides (PyMeI) strongly excite the brain and parelyse the extremities

² The potash can be greatly conomised, with a loss of some of the higher homologues, by rendating the distillate acid with hydrochloric acid, and concentrating it to a small bulk by evaporation at a gentile heat before adding caustio potash.

² Graville Williams destroys aniline and its homologues by heating with potassium nitrite and hydrochlone end. Hausermann converts the aniline into sulphate, which salt is much loss soluble than the sulphates of the other bases.

again treated with solid caustic potash. The layer of bases is ramoved, and further treated with stick potash or soda for several days, or until no more alkalı dissolves. It is only by prolonged contact with solid caustic alkali that the bases can be freed from water, and it is absolutely necessary to obtain them in a perfectly anhydrous state before attempting to separate them by fractional distillation This is a very tedious operation, but is greatly facilitated by operating in a vacuum, and by the employment of a Hemnel's tube or Henninger's or Glynsky's bulbs (Vol. I. page 14; Vol II 501) Goldschmidt and Constam (Jour Soc. Chem. Ind., 11, 159) found that the mixture of bases extracted by vitriol from coal-tar boiled between 92° and 200°, and after repeated fractionation a little passed over below 100°, and about onehalf between 114° and 117° (pyridine), while above this temperature no constant boiling-point was observed. Very little distilled above 160°. The most volatile fraction boiled constantly at 92°-93°, and was found to be a definite hydrate of pyridino, from which treatment with solid caustic potash caused a separation of absolute pyridine, boiling at 114°-115°.

C. Hausermann has pointed out that the amount of sulphure and employed in English the-works for treating 50 and 90 per cont benzols is insufficient to remove the bases. He found up to 0.10 per cent of prindine in commercial 50 per cent benzol, and 0.25 per cont in the toluol made from this. Hence the nearly pure benzone, toluene, xylene, &c, now largely manufactured, can be employed with advantage for the preparation of the printine bases, as the tedious fractionation has already been accomplished. Thus the base oxtracted by duited sulphure and from toluene will be nearly pure pyridine; from xylene, chiefly picoline, and from burning and solvent naphtha, the higher homologues English-made toluene yields about 0.5 per cent of pyridine, and a similar amount of picoline can be extracted from commercial xylene Pyridine is more commonly made from crude heavy naphtha, and procline from the highter ercoscote oils

Pyridine.

$$C_6H_6N$$
, or CH $\left\{\begin{array}{c} CH \cdot CH : \\ CH \cdot CH \end{array}\right\}N$

This body is the lowest and most important momber of the pyrdine series of bases. It has been used as an antasephs and germinde, and is employed in Germany for "densitivating" alcohol. Pyrdine is the starting-point in the proparation of several rehabile antipyroists, and many of the natural alkaloids are derivatives of it. The method of preparing pyridine from tare has already been sufficiently indicated. It may be obtained by several interesting synthetical reactions, as by passing a mixture of acetylene and hydrograme and through a red-hot tube $-2C_cH_a + \text{CHIN} = C_cH_b$. Pure pyridine is conveniently obtained in small quantity by distilling mootine and with lime $-C_cH_c\text{N} + \text{COOH} + \text{CaO} = C_cH_b\text{N} + \text{CaOO}$.

Commercial pyraduse may be purnfied by dissolving 200 cc in 400 cc (on a sufficiency) of strong hydrochloric and, filtering the figured in necessary, suit then adding 1000 cc of a 50 per cent. aqueous solution of potassium farrocyamide. The prespirate is a queous solution of potassium farrocyamide. The prespirate is aqueous solution of potassium farrocyamide of ammonia and the picolines are essally solutible, while the corresponding sail of pyradine dissolves but sparingly. The washing precipitate is treated with a cold, highly concentrated solution of caustic soda, when the pyradine separates as an oily layer, and, thus obtained, it contains a considerable but variable proportion of water, but if desired may be rendered anhydrous by treatment with stacks of caustic potash or soda, which should be innewed until they cosse to bunefy on standing.

Pure pyridine is a colouriess injuid, having a most powerful and noriestant odour, and producing a little taste in the month and at the back of the throat. The vapour causes severe headcade. Pyride has a specific gravity of 9858 at 0°C, and boths at 116.77 ascording to Anderson, or 115° according to Themus The presence of water, which it is difficult to separate completely and which pyridine absorbs with availty from the au, greatly reduces the boiling-point. Pyridine seems to form a definite hypothemic C.H.N. 3H,O. of specific gravity 1 0219, boiling constantly at 92°-93° C

Pynidme dissolves in water in all proportions, but is precipitated from its solutions by axcess of strong potash or soda. It is also miscible with alcohol, ether, chloroform, benzene, and the fatty oils. The effects of pynidine on animals are described on page 98.

Pyridine is a powerful base, neutralising acids completely and fuming like ammonia in presence of hydrochloric acid and other volatile acids. It blackens calomel, and precipitates many metallic solutions. Pyridine has no effect on a solution of calcum chloride.

¹ Pyridine might probably be advantageously purified from pyridi and strong-smelling impurities by dissolving it in petioleum spuri and passing hydrochloride acid gas, the preceptated hydrochloride of pyridine being lemoved, pressol, and dried at a gentle heat.

² According to A Ladenberg (Ber, xx; 299), the specific gravity of pyridine prepared from the mercuro-chloride is 1 0037 at 0° C.

but on passing earbon dioxide through the liquid calcium carbonate as prescriptated (No precipitate is produced if amiline be substituted for pyridine in this reaction). Absolute pyridine has no action on litrium, but in presence of water it turns it strongly blue, though the reaction is not capable of being employed for thinking the base, for which purpose methyl-orange is suitable. On phenolphthalem pyridine has no action.

Pyridine is an extremely stable body. It is unaffected by treatment with chromic or furning nitric acid, and these reagents may be employed to free it from aniline and empyreumatic impurities

When chlorus is passed into a chloroformic solution of pyradime, an additive-compound, C_{IT}N/C_S, separates in white flakes. Bromine forms a similar unstable compound. A substitutionproduct, d. br on pyrid in 6, C_{IT}H_SN_R, is formed by thesingto 200° a mixture of pyridine hydrochloride and bromine, or the onage-colured prespitate formed on adding promine to a solution of pyridine hydrochloride. It is prespitated by adding water to its solution in strong hydrochloror each, in needless melling at 100° but commencing to sublime at 100°. It is soluble in either and unacted on by akalles, acids, or existing agentless.

By reduction with tin and hydrochloric acid, pyridine is converted into piperidine, C₆H₁₁N, identical with the substance obtained by hydrolysis of piperine, the alkaloid of peoper

Dipy $i \& m_0 \in C_0H_{10}N_0$ so obtained with other products by heating pyridine with socium. Dipyridine is a base, which meltes the sublimes at a higher temperature in long needles, and forms a hydrocal horid a C_0H_{10} , M_0 HU), the solution of which pields with potassaum ferroquands a blue precapitate which dissolves in hot water to form a purple solution 1

Para-dippyridyl, C_bH_aN NC_bH_b formed simultaneously with dipyridine, is a base, crystallising in long needles malting at 114° and boiling at 305° (Jour Ohem Soc., xiv 483) Both these bodies yield iso-niotinios and on oxidation, while the isomen meta-dipyridyl yields in totinios and

SALTS OF PYRIDINA.

Pyridine forms well-defined salts, most of which are crystallisable and delaquescent. They are odourless when pure, and can be drad without change at 100°, but become slightly coloured on excessive to are and light.

¹ hos disperdane, Cu₂H_mO₂, as obtained by fractionating the mother-liquors from the preparation of dipyridine, is a yellow oil which does not solidify in a mixture of snow and saft, oven on addition of ciyatals of pyridine. It has a specific gravity of 1 08, and is a strong base, sparingly soluble in water, but missible mill proportions with alcohol and eltin.

Pyridine Nitrate, C₆H₅N,HNO₂, forms slender, colourless needles, or short thick prisms, very easily soluble in water, but less so in alcohol, and usoluble in ether

Pyridine Sulphate, (C₅H₅N)₉, H₂SO₄, is crystalline, and extremely soluble in water and alcohol ¹

Py usine Hydrochlorais, C₂H_NNHCI When pyrdune is neutral used with hydrochlora scul, and the solution evaporated at 100°, a syrup liquid is obtained, which, on coling, becomes gradually converted into a moise fraultaing crystals. The said deluçueses in moist au, and sublimes unchanged at a high temperature. It is notative of the converted into a very notable service that at 100°, and hence cannot be direct at the temperature without loss. It is readily soluble in water and alcohol, but mosobilor in either.

With platime chlords, a solution of pyridine hydrochlords, pields a yields or systalline preceptate of the chloroplatinate, (C₂H₂N;HCl)₂PtCl₄, essely soluble in boining water, less so in alcebol, and insoluble in ether. When gyridine chloroplatinate, free from excess of platime chlorode, as boiled with water for many hours, it is converted into the hydrochlorode of Pintino-Pyridine, C₂H₂PtN₂HCl₄(M), with blustion of 2HCl. Then substance is a sulphim-yellow, insoluble body, which evelves pyridine when boiled with caustic alless.

Pyridine Picrate, C,H,N,HC,H,(NO,),O, is deposited in beautiful yellow needles when picric acid in aqueous solution is added to a solution of an equivalent weight of pyridine. The salt has a remarkable tendoncy to carry pieric acid down with it, so that if twice the equivalent proportion of picuc acid be employed, the product has the percentage composition of an acid salt. Pv.2Pc: but its real nature is indicated by its behaviour with ether, which dissolves out the free picric acid, leaving the normal picrate Pyridine pierate may also be prepared by mixing strong solutions of sedium picrate and pyridine hydrochloride. The salt melts at 162° C., and is soluble in 91 parts of cold water, but in less than 6 parts of boiling water It is readily soluble in hot alcohol, but requires about 100 parts of the cold solvent, and is deposited on cooling in long, slender, interlaced needles of a beautiful vellow colour. It is only very slightly soluble in ether, chloroform, or benzene, and practically insoluble in petroleum spirit, but it dissolves with great facility in pyridine and cresylic acid. It is readily soluble on warming in ether, benzene, or petroleum spirit containing 10 per

¹ In Watts' Dictionary, vol. 1 page 755, there is only described an acul sulphate, which is said to be obtained by evaporating sulpharic acid with excess of produce.

cent of cresylic acid, and is freely soluble in aqueous solution of pyridine and sodium cresylate (A. H. Allen)

Pyridine picrate has an intensely bitter taste and nauseous pyridic after-taste. A moderate dose, for example 0.2 gramme, produces violent vomiting. It is a valuable insecticide.

Pyridine is remarkable for its tendency to form compounds with metallic salts These bodies are more or less hable to decomposition by washing or boiling with water, and lose pyridine when heated to 100°, or a somewhat higher temperature. The zinc chloride compound, ZnCl., 2C.H., N, separates as a voluminous white precipitate on treating an aqueous solution of zinc chloride with excess of pyridine. It crystallises from water in long. white silky needles, which, when repeatedly washed, or holled with water, decompose into pyridine and a basic zinc chloride. The zine chloride compound dissolves in hydrochloric acid to form a double chloride of zinc and pyridine, ZnCla (C.H.N.HCl)a, which forms groups of white lustrous needles Cadmium chloride behaves with pyridine in a manner similar to zinc chloride, the compound formed, CdCl, 2C, H,N, crystallising in needles and heing partially decomposed by a large quantity of water. The enpric chloride compound is precipitated in fine greenish silky needles on adding pyridine to an alcoholic solution of cupric chloride It is soluble in pyridine, in aqueous solutions of pyridine. and in ammonia. With mercuric chloride, a very dilute aqueous solution of pyridine (1-1000) yields a precipitate which dissolves extremely easily in warm water, and senarates out, as the solution cools, m long white needles. With mercuric 10 dide. pyridine forms a compound which crystallises from alcohol in beautiful white needles.

From acid solutions of pyridine, phosphotungstic acid throws down a very difficultly soluble precipitate

DETECTION AND DETERMINATION OF PYRIDING

The recognition and determination of pyridine are to a great extent based on the proporties and roactions sheady described. In the free state, the smell and basic character of pyridine amply suffice for its recognition in the absence of other base estabatences of powerful edour, and it is readily liberated from its salts by addition of caustics soda, and obtained free from every interfering substance by distilling its aqueous solution. It may also be extracted from its aqueous solution is made and the state of the state of the state of the cause soda.

In the absence of ammonia, or other bases, free pyudine may be determined by titration with standard and methyl-orange (not

htmus) i c.c. of normal acid neutralises 0.079 gramme of pyridine.

From antitine, pyridine is distinguished by not giving any coloured product on adding a solution of bleaching powder, though the liquid acquires a new and peculiar odour.

The presence of ammonium pyridine can be recognised (in the absence of fixed alkalies) by the red coloration produced in the aqueous solution by phenol-phithalem, on which pure pyridine has no action. If the indicator be used in considerable quantity, and a low temperature employed (as recommonded by J. H. Long, Analyst, xv. 53), the ammonis can be approximately determined by titating the aqueous solution with standard and.

K E. Schulze recommends ferric chloride as an indicator (see page 106). According to W. Lang, the traces of pyridine sometimes contained in commercial alcohol may be detected and removed by shaking the spirit with powdered zinc chloride, or, according to W. Kirsechmann, by the addition of an each solution of aluminium spliphate. In the former case, the pyridine is removed in the form of its zinc chloride compound, and in the latter case pyridine alum is formed.

The traces of pyridine sometimes present in fusel oil may be detected by adding pieric acid, which occasions a formation of pyridine morate.

"For the detection of traces of pyridine in commercial aminoma, H. Os is recommend that his sample should be nearly neutralised, when the odour of pyridine may be recognised. By distilling the nearly neutralised liquid, collecting the distillate in hydrochlorn scale, evaporating, and extancting the results with absolute alcohol, a solution is obtained containing but little ammonium chloride, as obtained to be able to the second policy of the containing the second policy of the containing the second policy of the containing the collection of the containing the collection of the coll

Commentar Preniums, as now produced, consets chiefly of pyridine and piceline Ammenia is apt to be present in notable quantity, as also pyriol and other strong smelling impurities A considerable but variable proportion of water is present.

Pyridine is employed in Germany, in conjunction with wood

¹ The pyridue produced at certain works becomes turbid when dilated with more than 40 per cent of water, whereas the best makes are muscible with water in all proportions. On distilling the former brands the disturbing impurity is left in the "tailing". spirit and turpentine, for "denaturating" spirit. An article intended to be used for this purpose is required to answer to the following official tests.

1. The colour must not be deeper than straw-yellow. 2. If 1 cc of the sample be dissolved in 250 cc of distilled water, and 20 c.c. of the resultant solution be treated with a 5 per cent aqueous solution of cadmium chloride, a distinct turbidity should appear in a few moments.1 3 When 100 c.c. of the sample is distrilled (in a small metal flask provided at the top with a small globe, which is connected with a Liebig's condenser, a thermometer being fitted to the globe, and a moderate heat applied) so that the distillate passes over in separate drops, 90 per cent, should have distilled when the thermometer stands at 140° C. 4. When the sample is mixed with twice its measure of water it must wholly dissolve, and no only drops must separate even after long standing. 5 Four drops of the sample heated on platmum foil over a Bunsen burner should burn with a sooty flame, and leave no residue. 6. When 20 c.c. of the sample is shaken with an equal measure of a solution of caustic soda of 1 4 specific gravity, a layer of anhydrous bases, measuring at least 18 cc (= 90 per cent), should separate out on standing.

The last test is now usually replaced by one prescribing the use of solid caustic potach. BO oc. measure of the sample is placed in a graduated cylinder, furnished with a stopper, and a long stake of potach immersed in it. The allast gradually absolo is the water from the pyridine, and forms a lower layer of saturated solution. A second stake is added as soon as the first has sunk much below the suiface of the pyridine, and is followed by a third if the second flugides completely or considerably. Agatacian should be avoided, and care must be taken that the last stack is left in contact with the upper layer of bases until the action is at an end. It is then cautiously removed with a bent wire, or broken down by a glass ord, and the volume of the layer of analyticus bases carefully observed. By this test, commercial pyridine smally shows from 8 to 10 per cent of water (= 92 to 90 ber cent, of anithrous bases).

Instead of determining the water, K E Schulze recommends titration of the bases with standard and. For this purpose 5 cof the sample should be dissolved in water, and the solution diluted to 100 cc. To 20 cc of this solution (=1 cc of the sample) as added 1 cc of a 5 pre cent aqueous solution of ferror chlorde,

We apper and Liders (Jour Soc. Ohem. End., vn 762) have pointed out the unreliable character of this test, which they attribute to the varying composition of cadmium chloride. Of two samples of the salt, only one gave the reaction with pyridne. They recommend the employment of a stronger solution of the pyridne than that prescribed in the test.

Normal sulphure and is then run in slowly with agitation, till the precipitated ferric hydroxide is redissolved 1 cc. of normal and (containing 49 grainnes of H₂SO₄ per litre) corresponds to 079 gramme of pure anhydrous pyridine, or to 095 gramme of pure anhydrous pyridine, or to 095 gramme of pure

Pyridine intended for pharmaceutical or medicinal use should not be altered by light; a 10 per cent. solution in water should not be reddened by phenol-philalein (presence of aminomia); and 5 cs, to which 2 drops of decinormal permanganate have been added, should return a red colour for at least an hour.

Piperidine. $C_8H_{11}N = C_8H_8(H_8)NH$.

This body has the constitution of a pyridine hexahydride. It is obtained by the reduction of pyridine by nescent hydrogen. The following formulæ show the relation of pyridine to piperidine and piperazine.

Piperidine is also obtained by rapidly heating pentamethylene-diamine (amylene-diamine) hydrochloride —

$$C_5H_{10}(NH_2)_2$$
, $HCl = C_5H_{11}N + NH_4Cl$.

Piperdune is also produced by the hydrolysis of piperine, $C_{18}H_{18}NO_9$, the alkaloid of pepper, which, on boiling with alkalies, splits into piperidine and piperic acid 3—

$$C_{17}H_{19}NO_3 + H_2O = C_6H_{11}N + C_{12}H_{10}O_4$$
.

Pipendine is a colourless impid iquid, of peculiar odour, resembling at the same time that of pepper and ammonia, and has

¹ Pyridine di and tetra-hydrides and then homologues are capable of existing theoretically The latter class, called piperides, have been prepared by the action of caustic sode and bromine on the piperidiaes (Ber., xx. 1645)

FITERALINE OF PIPERALIDER IS probably identical with diet by lensed at manne. It is a strong been seeling at 104-107, boing at 135-13, and absorbing carbon decords from the arr. Piperanne has nother counts one took properties, and passes through the system unchanged, but dissolves runs and in large amount, forming the neutral unste, QHan,Co,HAN,O, Piperanne phosphase forms four-acid tabular cystals, which channels dissolved the business of the business-order distinguish piperanne from a pen in ine, QHAN, or QHAN, or

⁸ A small quantity of piperdino is said to be obtained on distilling pepper with water alone, probably owing to partial decomposition of the piperine by water or a ferment (W. Johnstone, Analyst, xix, 46)

a very caustic taste. It boils and distils unchanged at 106°, and dissolves in all proportions in water and alcohol. When piperidine is treated with water heat is avolved.

Pspandine is a powerful base. Its aqueous solution restores the blue colour of reddened litmus-paper, and behaves like ammonia with metallic solutions, except that the precipitates produced with salts of zinc and copper are not soluble in excess. Pipendine shows earbon dixorde from the aux, and if the gass be passed into a solution of calcium chloride, to which pipendine has been added, calcium carbonate is precipitated. Piperdine may be estimated by titration with standard and, using either litmus or methyl-orange as an induster.

Piperdine forms a series of readily crystallisable salts, most of which are soluble The chloroplatmats, $(C_5H_{11}N)_2H_2PtCl_6$, forms orange needles, very soluble in water, but less so in alcohol

Pipersidine is a secondary amine. Whendropped into cooled methyl todde it forms the compound $C_{\rm ph}(CH_0)N,HI$. When distilled with alkalt his yields the free base methyl pipersidine, inc, which, when heated under pressure with methyl todde, gives the todde of dimethyl-pipersidine, the condition of the methyl todde, gives the todde of dimethyl-pipersidine, there are men pine. O.H. $C(H_0,N,II.$

The homologues of piperidine are called by Lade n burg pipecolines, $C_cH_{10}(CH_s)N$, inpetidines, $C_cH_{2}(CH_s)_2N$, copellidines, $C_cH_{2}(CH_s)_2N$, &c.

Piperuline is closely related to a number of the natural alkaloids besides piperine, as will be seen from the following formulæ.—

CONING. Dextro-
$$\alpha$$
-normal-
propyl-piperidine.

CH₂ $\left\{ \begin{array}{c} CH_2 CH(C_2H_7) \\ CH_2 CH_2 CH_2 \end{array} \right\}$ NH

Convergence Probable by (CH CH CH CH)

$$\begin{array}{c} \text{Consydenine} \\ \text{droxy-comme} \end{array} \text{ Probably hy-} \left. \right\} \text{ } \text{CH}_2 \left\{ \begin{array}{c} \text{CH}_2 \text{ CH}(\text{OH}) \\ \text{CH}_2 \text{ CH}_2 \end{array} \right\} \text{N}(\text{C}_3 \text{H}_7) \end{array}$$

Homologues of Pyridine.

The homologues of pyridine occur with that base in the products of the distillation of bones, coal, &c. Various members of the class have been obtained synthetically

PICOLINES METHYL-PYRIDINES. C.H. ; or C.H. (CH.)N.

Three isomeric modifications of picoline exist, differing according to the crientation of the CH, group in relation to the N. The pico-

line of coal-tar is chiefly the oitho-modification (1 2), often called a-nicoline, mixed with some meta- or B-nicoline (1.3)1 Although the former boils at 134° (Wendel; 129°-130°, Lange), and the latter at 140°, they cannot be separated by fractional distillation, but may be isolated by taking advantage of the different solubilities of their chloroplatinates (Ber. xn 2008) Lange (Ber , xviii 3436) thinks that a-picoline is preferably senerated from bone-oil by means of its sparingly soluble mercure-chloride, Its specific gravity at 0°, compared with water at 4°, is stated to be 0.9656 The platinochloride melts at 178°, the mercurochloride at 167°, and the picrate at 165°. The two last salts are moderately soluble in water. v-picoline (1.4) is produced by the distillation of acrolem-ammonia, or by heating allyl tribromide with ammonia, and by the reaction of pyridine on methyl iodide Its presence has been recognised in coal-tar, y-picoline is stated by A. Ladenburg (Be., xxi. 285) to boil at 142° 5-144° 5. the specific gravity being 0 9742 at 0° C. The platinochloride melts with decomposition at 231°, the aurochloride at 205°, the mer-curo-chloride at 128°-129°, and the picrate at 167° These characters are not strictly in accordance with the observations of Lange (Ber . xvni. 3436)

The precluses are metamers with a nilnne, CoH, NIL, which, however, is a primary arms, whensa the prolines have the characters of textary bases. In their odour, solubility, basic properties, and characters of their salls, the proclines closely resemble the lower homologue pyridme, but have a lower density and higher bothner-coint than the latter body.

LUTIDINES C.H.N.

The bases of this formula may have the constitution of ethylpyridines, $C_5H_4(C_2H_5)N$, or of dimethyl-pyridines, $C_6H_4(C_2H_5)N$.

- 14 or y-ethyl-pyridine constitutes the greater part of coal-tar lutidine. It is a colourless high of 9443 specific gravity at 0°, boiling at 154°, and miscible with cold water in all proportions. By exidation it yields is o-nicotinic and
- 1. A. Lad on burg (Rev., vatir 2888) affirms the emission of two 8 pieces, the variety from glycord boling at 1476-147 (uncontexts), and that from strybhine at 146-147 (uncontexts). C. 8 to a kr (Rev., xux 1311) depicted and the stress of the stress o

A β -ethyl-pyndine is formed, together with its lower homologues, by heating glycerol with acetamide and phosphoric anhigh dide (C. Stoe hr, Jour. Frac. Ohem. [3], xhm. 163). It boils at 140²-146², has a specific gravity at ${}^{4}/_{1}$ 0 or '9761, is almost insoluble in water, and realth snotwine and on oxidation.

Three moments of m et hyl-pyrid in es have been found by Recemberg in vitrol-tar Of these, left 1.2 : 6($-\alpha_c$) isomeries as a colourless oil boiling at 142°-143°, and having a penetrative odour resembling that of oil of peppermint. It is freely soluble in cold, but less so in hot water. The 1.2 · 4 (α - γ) isomer boils at 167° The 1.2 : 3 (α - γ) modification has not been isolated, but its presence is inferred from its product of oxidation, 180-110 clumber.

Hanseh (Annales, coxv. 1) has described a lubdime $(O_{21}G(\Omega_B)_{1})$ bohme at 164°, obbaned by distilling a muture of lubdine-trearboxylate with lime. A lubdine, apparantly having the constitution ββ-dimethyl-grudine, has been prepared by Durkopf and Gottsch (Ber, xmi 1113) by simunating O_2 from a dimethyl-grudine-carboxylic and obtained by the condation of a parv of line boling at 160°-217°. It boils at 169°-170°, has a feeble, not unpleasant odout, and dissolves sparingly in cold, but readily in boling water. The specific gravity at $θ_4$ is 0 9614. The measure-chloride crystallises in long sparingly soluble needles, melting at 170°. On condation it yields a pyridine-dicarboxylic and melting at 314°-315°, from which fact, and its external characters, the authors infert it to be d into cotinite a ctd.

COLLIDINES C₈H_UN

A Hanzsch Amalen, cexv 1; Jose Ohem. Soc, xliv 82) gives the following description of the two known modifications of collidine.—

	a Collidme Mathyl-ethyl-pyridme C ₅ H ₃ (OH ₂)(C ₂ H ₅)N	β -Collidane β -taimethyl-pyrkline $C_5H_2(CH_3)_5N$
Boiling-point,	178*	171*
Specific gravity at 15°,	853	917
Solubility in water, Behaviour on exposure to air.	Very slight Unchanged	More readily soluble in cold than hos Becomes brown
CgHijN, MAuCl ₄ , Addition of CrO ₂ gives Mn, Co, and Fe salte, AgNO ₃ ,	Does not melt under water Red oil No precipitate, No precipitate	Melts under hot water, the dry sall melts at 112 dry sall melts at 12 dry sall met 12 dry sall melts at 12 dry sall met 12 dry sall melts at 12 dry sall met 12 dry

O. de Coninck has described a \$\textit{G-collidata} bolling at 195^-.

196^* (Compt. Rowl, xur 296, xvr. 298), having a specific gravity of 9556 at \$0^*\$, and another modification, stated to be a trimethylyridine, has been isolated by \$J\$. Mo \$h\$ is \$(B\textit{G-g}, xxx)\$ 1006, \$J\textit{Gomes.} Soc. http://dx. 250. http://dx. 250.

Pyridine-Carboxylic Acids.

Fyrnine itself is an extremely stable body, assuting the strongest oxidising agents, but its homologues yield by oxidation a series of axidising agents, but its homologues yield by oxidation a series of such in which the allyl-groups are replaced by a corresponding number of carboxyl-groups. The pyrnine-carboxylic acid derive their chief interest from the light they throw on the relationship of the natural vegetable alkalouis to the pyrnine-bases. Three sections principles and acid may be a consequently and the strong part of the natural vegetable alkalouis to the hyrnine-bases. Three sections of piculia can one and acid and acid control of the strong of the strong

PYRIDINE-MONOGARBOXYLIO Acids, C₅H₄N COOH,² umbe in themselves the basic cheracters of pyridine with those of an acid. Thus they combine with hydrochloric acid, and the resulting com-

1 The pyridine monocalboxylio acids have the empirical formula $C_0H_2NO_2$, and the same percentage composition as nitrobenizens

The bases from coal-tar bolang between 180° and 140° are bolled in an apparetas furnished with a serlius condense with the times than weight of possum permanganates in 2½ per cent, augmons solution, until the permanganates in 2½ per cent, augmons solution, until the permanganates in 2½ per cent. The permanganates is 22 and 12 and the clear liquid concentrated to a small bulk. It is then mentialed and treated with sociator obegier. The prespirates separated, decomposed by subplimited hydrogon, and the ditrute decolorised by animal charcost. On further consensation and cooling it deposits colonious mealing of pros 1 ni a sold. The district from the cooping precipitate in further evaporated, another desired and the coloring point with occupied of pros 1 ni a resulting blushingeron precipitate in the permanel, belief and point with occupied by sold in public prospers of the permanel of the permanel p

pound forms double salts with mercune chloride, platinic chloride, &c., while, on the other hand, they form a series of well-defined crystallisable salts. The following table exhibits their more important characters:—

	Or the Compound Or a Acid Puolinia Acid	Meta Compound or 6 Acid Nicotento Acid	Para-Compound 01 y Actd Itonicotimis Acid
Mode of formation, .	Oxidation of a- picoline by per- manganate	Oxidation of \$\beta\$ picoline by permanganate, or nicoline by permanganate chromic acid or nitricacid	Action of heat on pyriding di- or tri - carboxylio acid Oxidation of y physika
Crystalline character,	Prismatic needlee	Needles	Needles.
Melting point,	185°, sublimes in lustrous needles	229*-221*	806* (299*) (209*), sublimes in tab- ular crystals
Solubility,	Restly soluble in colden hot water and in alcohol Nearly insoluble in ether, chloro- form, benzene, &c	Sparingly soluble in cold, easily in warm water, aparingly in either or obloro form	in water, very sparingly in
Reaction with nentral lead acetate,	No change	No change,	
Reaction with am monlacel lead acetate,	No change	White orystalline precipitate.	
Reaction with cupric acetate,	Slowly deposits shining lamine and needles of violatiblia col- our, and metal- lic instre Sci- uble in hot water	precipitate, in- soluble in a large quantity of water,	Green precipitate on warming
Reaction with ferrous sulphate,	Pale reddish yel- low coloration	No change	No change
Cheracters of hydro chloride— CaHsNO2,HO1,	Large, lustrous, ortho-rhembto prisms, which become rapidly turbid on ex- posure to air.	ouste permanent	Large shining crys tals.

On heating with lime, the above acids yield pyridine, just as benzore and yields benzore under similar conditions. The sodium salts of the α and β acids, when treated in solution with sodium amalgam, give off ammonia, and yield the salt of an unsaturated and of the fatty sories, $C_{\rm sL}(Q_{\rm s})$.

Pyriding-dioarboxylio Agids. $C_6H_2(COOH)_2$ Of the six possible acids of this formula, all are known. They are produced by the exidation of homologues of pyridine contaming two

substituted hydrogen atoms, and also by the oxidation of other substances

Quanolatic Acta [the a-S modification] so obtained by the oxidation of coal-tar quinoline by permangemate, and is the analogue of phthalic said, obtained similarly by the oxidation of naphthalices. It crystallness in short prisms, slightly soluble in cold water, more readily in hot water and sloobe, insoluble in beatzene. It blackens when heated, and molts at about 228°, apparently being converted into monthine said (Jose Chem Soc, xiv. 90). The acid is issurored from; its equous solution by ether.

Latidine Acid [a-y] is similarly produced by the action of permanganate on unohomne-quinoline It melts at 235° (219'), forming sensitions acid, is spanngly soluble in old water, and gives with cupric acetate a pale blue procupitate. (See Berichte, XX. 137')

Dimeolinic Acid [a-a'] melts at 226°; Isocinchomeronic Acid

[α-β] at 236°; and Dinicotine Acid [β-β] at 323°.

Conchanaeomic Acad $[\beta \rightarrow \gamma]$ as the chaef product of the oxidation of quantum by nature such and as she obtained, together with other products, by the similar treatment of cinchanne. It expatallass are white pramatic needles, which made at 25% (267%) with partial decomposition, and is only very sparnagly soluble, even in bothing water. It forms two classes of salls I to most characteristic reaction is its behaviour with cupius costas, which does not give a precapitate in the cold, but on heating the bignid becomes turbid, clearing again on cooling. On prolonged bothing, a nermanent assum-blue representate is formed.

All the dicarboxylic acids which contain a carboxyl-group in the a-position give a reddish yellow coloration with ferrous sulphate

Principute-rancarocture Aones, C.H.(CO.OH), are obtained by the extraktion of certain alkaloids. Thus quinne, quinding, and caneboundune, by beiling with an alkaline solution of permanganana, yald high organizations are acid, which forms orthorhombus princes molting (with blackening) at 944°, while betherms, when excludiby nitre each, yalds in isomorne body be thorous and, crystaliing in the trichine system. Both acids give a deep red colour with ferrous sulphasis, destroyed by a mineral acid.

Pyridine-Tetracarsoxylat Acids, C₅HN(COOH), have been obtained

PTRINISS-PRYMARSONIZED AGE, C.N.(COOH), forms crystals containing 2 ages. It becomes anhydrous at 130°, and decomposes without melting at 230°. It is freely soluble in water, and is a strong acd, resembling oxide and in its power of form gard and double salts (Hanssch, Jour. Ohm. Soc., XIV. 55).

PYRROL 113

Pyrrol. 1 CaH, N, or CaH, NH

This associate of the pyridine bases I is a colourless liquid of pungent taste, and odour like that of chloroform. The specific gravity is 1077, and boiling-point 130°–133°. It is but tittle soluble in water, and nasoluble in alkalies, but dissolves in dilute cackla, shoolid, and other. It is indifferent to most respents, but appears to possess feebly—marked bases properties. The only definite sait is the normate, which forms untable red needless melting at 71°.

Pyrrol turns brown in the air, and when warmed with acids forms a red substance known as pyrrol-red, the reaction apparently occurring being $-3C_1H_NN + H_0 = C_{12}H_1/N_0 + NH_3$. A piece of pine-wood, moistened with hydrochloric acid and exposed to the vapour of pyrrol, becomes deep red

to the vapour of pyrrot, becomes deep red

When a cold aqueous solution of seatm is treated with pyriol and a little dulute sulphure cad, a heavy blue precipitate, resembling indigo, is obtained. When both resgents are dissolved in glacula sectic acid and boiled, a deep blue solution is obtained, apparently containing the same colorung-matter.

If a solution of phonanthrane-quinone in sects and be tested with pyriol and a little dilute sulphune and, a brown presuptate is formed, which dissolves in chlosform with a beautiful violet adoious. When an aqueous solution of bearo-quinone is tested with pyriol and dilute sulphune acid, a dark green precipitate is formed, insoluble in other These reactions indicate this close relationship between pyriol and thin oph ena, which itself has the constitution of a thin of fur fur an Many of the reactions of pyriol are also produced by car baso, which is an infid-dip henyl. Indofe has a constitution between pyriol and carbasol. Thus —

CH·CH. CH CH NH	$\left\{ \begin{smallmatrix} Indols, C_0H_7N \\ CH & CH \\ .C_6H_4 \end{smallmatrix} \right\} NH$	$\left\{ \begin{array}{l} \text{Carbanol, } C_{12}H_{0}N \\ C_{0}H_{4} \\ C_{0}H_{4} \end{array} \right\}NH$
CH CH O		$ \left\{ \begin{array}{c} C_0H_4\\C_0H_4 \end{array} \right\}O $
Thiophene, C ₄ H ₄ 9 CH CH S CH CH S	Thionaphthene, CaHeS! CH CH CaH. S	

Pyrrol has been obtained synthetically by passing acetylene and aminoma through a red-hot tube, and also by the dry distillation of the aminomium salts of muote and saccharic acids.

² The proportion of pyrrol contained in coal-tax is very small. It is best pregared by shaking bone-oil with dilute sniphuric acid and fractionating the insoluble portion. The fraction boiling between 100° and 150° is heated Vol. III PART II.

H

114 IODOL

Two somere methyl-pp role casts in bone-oil, besides a dimethylpprof, boning at 165°, which has also been obtained synthetically, and closely resembles pyrol. In the homologues of pyrol occurring in bone-oil, substitution has always occurred in the C_iH_i group, but by the action of alkyl addies on potassum-pyrol substitution of the hydrogen of the NHI group can be effected.

TERRADIO-PERSON, CALNII, has been seconify introduced into medicine under the name of "10 of 0." It is prepared by the ecton of sodreed potessum todie on pyrnol, and forms a tasteless, pale yellow, cytakline powdes, having a faut thymol-like oblight to such as part of the prepared in 100° int gives off action wapour at a somewhat higher temperature Iodol is nearly moduled in water, but readily in either and chloroform. It theselves in three parts of alcohol, and the solution is presquisted by adding water, but not by glycard Iodol contains 90 per cent of todine and possesses antisoptic am Iodol contains 90 per cent of rodine and possesses antisoptic move which it is alghi todour and freedom from toxic properties give it the preference Iodol can be necessated by the green colour of its solution in sulphure and, and by the figure along or its solution in sulphure and, and by the hight red colour produced when an alcoholic solution is warned with intra and

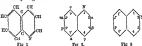
QUINOLINE AND ITS ALLIES.

The interesting base which gives its name to the quincline series bears the same relation to naphthalene that pyridine bears to bessens, that is, it is derived by the substitution of an atom of introgen for one of the CH groups of naphthalene (see foot-note, Vol. II page 507) —

with a large excess of solid canetic potash in a reflix apparatus until the whole is fused, when any unchanged oil is esperated and the crystalline mass of pota six um pyriol, Q_iI_kIKN, in powdered, and after being washed with other is treated with water and distilled with steam, when the pyriol is regenerated.

1 To notate these methylypyrots, the faction of hone-oil boling between 10°s and 150°s occurred in the poissame derawtee, and this is bested to 20°s in a steam of curiou dours! Two moment home pyriotic action 20°s in a carde as a formed. The a nach milet at 160°s, and form a lead acti vary comble in water, while the β and melts at 160°s, and form a lead acti vary somble in water, while the β and melts at 160°s, and form a marry monthle lead act in 10°s, and form the corresponding a- and β-homopyrrols are reguerated. The first boils at 145° and the latter at 148° at 743 mm, pressure

Quinoline may be represented by the following constitutional formule: Where substitution occurs in the profine-nucleus, a, β , and γ (or P-1, -2, and -3) products are obtained, while substitution in the beausen-nucleus pedies orthor, made-, power, and anadeurvatives (or B-1, -2, -3, -4), according to the position of the substitution λ by the profit of the substitution λ by the substitution λ between λ by the substitution λ



Just as two isomeron naphtholes exist, so two isomeric quinolines are theoretically possible, and appear to have been obtained. This the quinoline obtained by distilling quinne, emclouine, and other alkaloids with potats (fig 2) appears to differ in some of its reactions from the quinoline contained in coal-tai, which is often called it we old line (fig 3). On the other hand, Houge we rift and Van Dory (Jone Ones, Soc., xiv 89) contend that the quinolines obtained from both sources are adentical

A whole series of higher homologues are produced, together with quinoline, on distilling alkaloids with caustic potash. ', -medhyl quinoline or lepidine, C₃H₃(OH₂)N, the first member of the series, boils at 266° Of the next member, despotane, C₁H₁₂N, and the still inches homologues, very tittle is known.

A parallel series of bases have been found in coal-tar and shale oils. They are obtained from the fractions of the bases boiling above 200°, and hence distil after the pyraline bases have passed over Cuthaddrine, or a-me thyl qui nolline, C₆B₄(CH₂)N. bools at 239°, and sometimes forms 25 per cent of coal-tai quinoline. It is a colourless inquid (also obtainable synthetically), the condition of which yields either a benzane or a quinoline derivative, according to the nature of the oxidising agent * Irridohne, somerous with quindleine, and probably identical with lepidine, is also con-

If the distillation be conducted in presence of copper oxide, the quinoline obtained is almost free from higher homologues

• When quandities is bacted with anyl noble it forms the compound $C_0H_0(H_0/QH_0)H_1)$, which on heating with caustie potable a converted the α varning $\alpha_0H_0H_2$, $H_0(ago 118)$. A similar body is obtainable from leptimic, and a muttine of the two has been used for dyrang agh, to the colour very furtire. When bested with philable chilydrice, quandities resuct to form a body of this philable in class known as quantities qallow (see Vol. III Part I page 1/4V).

tained in coal-tar. It boils between 252° and 257°, and yields a crystallisable intrate, chromate, and hydrochloride.

From the acid tar produced in the purification of shale-oil, Robinson and Goodwin (Trans. Roy. Soc Edin, xxvin 561, xxix 265) obtained the following bases of the quinchine series

Basi	,		Formula	Boiling Point, * 0
Tetracoline,		-	C ₁₉ H ₁₈ N	200-205
Pentacohne,			C ₁₃ H ₁₅ N	805-810
Hexacoline,			CMH37N	825-880
Heptacoline,			C10H19N	345-350
Octacoline, .			C ₁₆ H ₂₁ N	800-866

Quinoline, Chinohne CaH,N.

This base is formed by distilling quinine, cinchonins, or strychnine with aqueous potash, and by other interesting reactions, but is best prepared by shaking together nitrobenzene (48 parts), aniline (76 parts), glycerin (240 parts), and sulphuric acid (200 parts). When the aniline sulphate has dissolved, a reflux condenser is fitted to the flask, which is heated to 130° till reaction sets in, when the flame is removed. In about three hours, or when action is at an end, the product is cautiously diluted with water, and boiled to get rid of traces of mitrobenzene, after which lime or caustic soda is added, and the quinoline and unchanged aniline distilled over in a current of steam. The oil obtained is separated from the acusous layer, dehydrated over caustic potash, and fractionally distilled, whereby a separation of the bases is effected tolerably readily, and an boiling at 184°, and quinoline at 239° To purify the latter it is again fractionally distilled, and boiled with weak chromic acid mixture (to oxidise any amiline), or the quinoline is dissolved in six parts of water, and strong sulphunc acid added in the exact quantity necessary to combine with the base. After cooling, the liquid is filtered, and the insoluble acid sulphate washed with alcohol till snow-white, and then decomposed by potash 1

1 The reaction in the foregoing reaction may be written thus .-

 $2C_8H_7N + C_8H_8NO_9 + 8C_8H_8O_9 = 3C_9H_7N + NH_9O$.

The change is undoubtedly due to the formation of acrolein, C₀H₂O, from the givenin, and the reaction of this with annihe to form acrolein-aniline, with simultaneous oxidation by the nitrobenzene —

$$C_9H_8O_3 + C_6H_5NH_2 + O = C_9H_7N + 4H_2O$$

The homologues of quinoline may be obtained in an analogous manner, and by

Quinoline is a colousless mobile biquid, having a posterating and peculiar taste, and an after-tasts slightly resembling pepperumit-oil. It has a faint aromatic odour, like that of bitler-dimond oil. Quinoline eviporates complietly but slowly at the oothumy temperature, so that the grease-spot formed by it on paper is not permanent. It boils at 285"—239", according to most observers, 231"5, according to S, pal & n let 2; and 241"3, according to K, retainly, like specific gravity is stated; to be 1081 at 0"C, and 1094 at 20°C, compared with water at the same temperature

Quinoline is very spaingly soluble in cold water, but more freely so in hot. It is miscible in all proportions with alcohol, other, carbon disulphide, and fixed and volatile oils, and is also easily soluble in chloroform, amylie alcohol, benzene and petroleum spirit.

On exposure to air, quinoline becomes resinified

Quinoline has well-marked basic characters, and forms an extensive series of salts, most of which are crystallisable and deliquescent. It precipitates ferric and aluminium solutions, and at a ligh temperative decomposes ammonium salts

Quinoline can be titrated fairly accurately with standard acid, if methyl-orange be employed as an indicator

REACTIONS OF QUINOLINE AND ITS SALTS,

Quinoline salts in aqueous solution are precipitated milky white by caustic alkaites and ammonia, the precipitate being squewhat soluble in excess. From the alkaline liquid, the quinoline can be readily extracted by either olloroform, or petroleum sourit

Iodissed nodide of potessium gives a reddish-brown presquisited vers in third solutions of quinoline salts (1 in 20,000) Potassomercuric nodide only precipitates quinoline from tolerably strong solutions (1 in 3000), the prespitate engine pellowsia white and amorphous, but converted into delicate ambor-yallow meedles on addition of hydrochloric said. This reaction is characteristic. Phosphomolybids acid, in presence of mirro said, produces a vallowsia-bridt presentate in unpoints solutions.

Potassium ferrooyanide colours solutions of quinoline salts reddish, and on addition of hydrochloric acid a reddish-yellow amorphous precipitate is thrown down, if the liquid be not too dilute

Quinoline is precipitated by piecic acid, but not by tannic acid or force chloride, and its salts, in the solid state, yield no colourreactions with nitric acid or strong sulphunc acid, either alone or in association with oxidising agents

With potassium bichromate, if carefully added, quinoline salts

employing derivatives of aniline or its homologues, quinoline substituted in the hencene-ring may be obtained

yield a precipitate of delicate dendritic crystals of the high romate $(C_9H_7N)H_9$, C_7Q_7 , said by Donath to be soluble in excess of the reagent. Quinoline bichromate melts at 165° C.

When quinoline is heated with sodium, diquinolyline, CyHeN.CyHeN, snalogous to dipyridyl and diphenyl, is formed the polymerised, quinoline yields yellow needles of diquinoline, (CyHeN).

When quankine and amyl todalo are holied together for a short time, they combine to form a crystalline body containing C_HH₁(C_HH₂)NI If the product be dissolved in boting water, and the solution filtered and bouled with causet sods or ammonia, avading access, a blue colouring matter is formed, which, can allowing the liquid to cool, is preceptated, leaving the solution and body cystallinging in green plates, having an establic dustre the nearly manifolds in cold water, but dissolves in alcohol to form a rich currently blue solution. Which deve all the line and the method of the colour colours and the colour colours and the colours are colours and the colours and the colours are colours.

The foregoing reaction, as also that with potassium bichromate, is said not to be obtainable with the quinoline (leucoline) of coal-tar

Quinoline possesses powerful anisophic properties 0 2 per cost, of the tartists is said to completely prevent the lacht formentation of milk, the decomposition of unne and gelstin, and the development of bacteria in cultivation-fluid. Even in contentated seliment if does not congulate abitantin, and in the proportion of I per cent it completely destroys the coagulability of the blood. On the other hand, quinoline is remarkably inactive to yeast-cells, and does not affect the alcoholic fermentation, even when present in considerable cumnity

Quandum has been used in medicine as an antipyretic, the adult does of the tarknets being from 7 to 12 grains. It is said by some not to produce any unpleasant after-effects, but by others to cause irritation of the stomach and collapse. It is not found in the urine of those who have taken it internally.

Commercial Quinchine is often very impure and quite unfit for machemia use C. Extn. (Pharm Jour, [3], xm. 661) has described a specimen which had a deep brown colour and in adour hise oil of faitter almonds. On treating at with hydrochlorio scale a large proportion remained insoluble, and was evidently unconrected mitrohensene, whale the soluble part gave the reactions of anilms.

Canchonine-quinoline often centains lepidine. Such samples give the cyanine reaction (see above) with amyl iodide and caustic alkali

The salts of quinoline should be completely soluble in water,

and the free base in a slight excess of hydrochloric acid. The neutral solution should be free from bitter taste (which indicates the presence of impurity derived from einchonume), and should not give a coloured preemitists with caustic alkalies.

Counciline Twi haté, (C,H_N), (C,H_Q), a now used extensively in medicine. It melts at 125° C, and possesses the advantage of being permanent in the air, whereas most of the salts of quincine are dalquescent. It this olives in 80 parts of cold water, in about 160 parts of rectified spiris, and in 360 parts of either. It produces much the same effects as sulphate of quining, and is given in similar doses, but is far lower in prose

Quantitine Hydrochloride, C₈H₇N,HCl, melts at 94° C, and sublimes unchanged It dissolves in water, alcohol, and chloroform, and sparingly in cold ether and benzene

Tetrahydroquinoline. $C_0H_{11}N$; or C_0H_4 $\begin{cases} CH_2 CH_2 \\ NH CH_2 \end{cases}$

when quinoline is select on by assent hydrogen, it is his converted into d in γ dro qu in oll in ζ , ζ , ξ , ξ , as said body melting at 161, and subsequently into total n γ dro qu in oll in α , which is a liquid bothing at 2465. Both these reduction-products yield introsminae, and can be alkylated, and hence are secondary bases. Tetrabydroquinoline possesses stronger unityretic athanteriers than quincline itself, and thus property is exhibited still more strongly in certain of its derivatives, several of which have received some application in medience (see below)

Antipyretics allied to Quinoline.

A considerable number of new substances related to quantine, and mostly alled to testabydroquantine, have been recently introduced as febritages and antipyretics. Some of these are very powerful in then action, and appear likely to receive a permanent place in mediteme, but they are not periodics, and cannot be substituded for quinne in cases of agee or internitient forers. The following are the most important of the antipyieties derived from or related to quinchine.

M-KAROLINE is the acid sulphate of a base having the constitu-

¹ Other archypyrelocs are described in the sections on ambides, smulophenoly, antipyrue, for Many interesting facts relating to and telationships of the antipyretics have been collated by T S Dymond and an anonymous German authon (Piners Jaur, 181, xwn 386-386). A fuller and more executed description of them agree nia series of studies on "Modern Mateua Medica," contributed by H. He I bing to the British and Colonial Druggies, 1891, and since published in a separate form.

tion of methyl-tetrahydro-quinoline, C₀H₁₀(CH)₈N, obtained by reducing quinoline by tin and hydrochloric acid, and reacting on the resulting tetrahydroquinoline with methyl include

A-Karoline had a similar constitution, but contained ethyi, Co-H_{st}, instead of the methyl-group

M-Karmer is the hydrochlorade of Hydroxy-tetrahydro-methyl-quinoline,

C₀H₃(OH) (CH₂CH₂ | The corresponding ethyl-derivative is known as A-Kań ine

On adding a caustic alkali to the aqueous solution of a kairine, the penetrating characteristic odour and bitter taste of the free base are easily recognised, while the alkaline solution rapidly becomes coloured and deposits a brown humus-like substance When the aqueous or alcoholic solution of a kairing is treated with an oxidising agent, such as potassium bichromate and an acid, it gives a series of colours ranging from violet-blue to purple, or sometimes greenish, Without the addition of an acid, the solution becomes dark purple, and on standing a violet precipitate is formed, which dissolves in alcohol with black colour. A drop of ferric chloride, added to a dilute and neutral solution of kaurine, instantly produces a violet coloration, rapidly changing to brown, with precipitation An excess of ferric chloride added to a strong solution of kairine produces a nearly black precipitate Sodium nitrite and dilute sulphurio acid produce an orange or red colour in kairine solutions. Potassium ferrocyanide gives a voluminous precipitate, and phosphotungstic acid a pale yellow precipitate

The kairines act as powerful antipyretics. Their use is almost obsolete, as their action is somewhat uncertain, and they are said to be liable to produce vomiting, cvanosis, and collaise

THALLING is the commercial name of another analyzetic, metameric with m-kairine, and having the constitution of a sait of tetrahydro-paraquinanisol ---

$$C_0H_2(O.CH_2) \cdot \left\{ \begin{array}{l} CH_2 CH_2 \\ NH CH_0 \end{array} \right\}$$

Thisline is prepared by heating paramido-minel and paramito-masel with givenn and subjunct each, and reducing the product with nascent hydrogen. Thisline base crystallises in large colour-less rusms, having a bitter, subne, and pumper tasts. It make at 42° C, and is spaningly soluble in water, but readily in alcohol, ether, chloreform, or benzeen

Thalline Sulphale, (C₁₀H₁₃NO)₂H₂SO₄ + 2H₃O, is the most common variety of commercial "thalline." It occurs as a yellowishwhite, granuler or crystalline powder, having a bitter, aromatic faste.

and a faint odour resembling anise and meadow-sweet. It dissolves in seven parts of cold water, but only sparingly in alcohol, and the solutions become darker on exposure to light. A very dilute aqueous solution of commercial thalline gives with ferric chloride a yellow coloration, changing to emerald-green (destroyed by reducing agents), and passing in a few hours to deep red. The reaction is extremely delicate. A green colour is also produced by auric chloride, argentic nitrate, mercuiic nitrate, chlorine-water, &c., and, in acid solution, also by solution of bleaching powder and potassium ferricyanide. Strong sulphuric acid dissolves thalline sulphate without coloration, but on addition of nitric acid the liquid becomes deep red, and immediately afterwards yellow-red Fuming nitric acid colours a dilute aqueous solution reddish. Sulphuric acid and sugar give a red coloration. Indine colours the solution dark brown, then dangy green Ammonia forms a white precapitate of the free base, readily taken up by ether on agritation. If not too dilute, solutions of thalline sulphate yield precipitates with the general reagents for alkaloids

If to an aqueous solution of β-naphthagumone a small quantity of the solution of a thalline sait be added, and then a drop or two of caustic sodia solution, a fine cherry-red coloration is produced, becoming more brillant on adding nitro acid. The colouting matter is extracted by either or chlorofoim

Thallune Tartrate occurs in commerce as a yellow-white crystalline powder I the dissolves in ten pats of cold water, and the solution gives the same reactions as the sulphate In alcohol it is very sparingly soluble The salt contains 52 2 per cent of thalline.

The salts of thalline become altered by exposure to light

Thaline salts are powerfully antipyretic, and have been employed in yallow favor. They cause profuse persuptation, and an ept to profuse depression, &c. Hence their internal use is practically obsolete. Thaline acts as a direct blood-posson, its antithermic proparties being due to the destruction of the real corpussless. It has found considerable application in the treatment of genorrhos. The sulphate is official in the German Pharmacopera of 1890.

Exhibition of thalline causes a dark colonation of the turne A derivative, which also gives a green colour with ferre chloride, but differs from thalline in being extracted by agitating the acidulated turne with petroleum spirit, should first be removed, and then the unallered portion of the thalline can be isolated by rendering the turne alkaline with ammonia, and agitating with either or benome. Very small quantities of thalline can in this way be reorganised in urine

ETHYL-THALLINE, CloH10ON(C2H5), is produced by heating ordinary thalline with ethyl iodule

THERMIFUGIN IS a name given to the sodium salt of methyltrihydroquinoline-carboxylic acid.—

$$(COON_a)C_0H_a: \left\{ \begin{array}{l} CH_2 CH_2 \\ N(CH_3).CH_2 \end{array} \right\}$$

Quinazolines.

By the replacement of one of the CH groups of quinoline by N, bodies are obtained which bear the same relationship to quinoline that the azines bear to pyridine Thus —

A substituted meta-quinazoline having the constitution of a phenyl-dihydrophenmiazine —

$$N \left\{ \begin{array}{c} CH N(C_6H_5) \\ C_2H \end{array} \right\} CH_2$$

has recently acquired some practical interest as the base of "0 r o x 1 n" a preparation said to have valuable tonic, stomachic, and appelating properties, on which, however, some doubt has been thrown (*Pharm. Jour*, [3], xx 709, 825, 977, xxi 43) The usual dose of orexm is from 2 to 10 grains

OREXIN, which occurs as a hydrochloride having the composition $C_{1,1}\Pi_{\mu}N_{\mu}HCI+2\Pi_{\mu}O$, is prepared by reacting on the sodium-derivative of formanilide by ortho-nitrobenzyl chloride, according to the equation—

Na(CHO) N C₀H₈+Cl CH₂ C₆H₄ NO₂-NaCl+CHO N(C₀H₂) CH₂ C₄E₄ NO₄

The mitrobenzyl-formanlide, on reduction with tin and hydrochlore and, forms the closed chain compound which is the base of orexm — OREXIN. 123

$$CHO N(C_0H_5)$$
 $CH_23H_2 = 3H_2O + N \{CH.N(C_0H_5)\}$ H_2

Occun (hydrochlorods) crystallises with \$H,\(0\) in white needles, melting a \$60° When kept under an excentor for some they become anhydrous, and then mult at \$21°. Orcun has a bitter task, and consenhant intens, burning after-tasks. The powder induces violent sneezing Orxun dissolves readily in water (13 parts) and alcohol, but not in other On adding an alkali to the aqueous solution the free base is separated as a winte floculant pre-upstate resetly shalble in other and chlorofaum. A solution of orxin yields with mercure chlorate a white precipitate solution that the second coloring to occur a coloring to occur and the coloring to the second coloring to the coloring to the second coloring to occur and the coloring to occur and the second coloring

On heating orexin in a test-tube with about twice its measure of nin-clust, the strong characteristic odour of phenyl-monitories produced. On treating the residue with hydrochloric eard, and adding bleaching-prowder solution to the filtered liquid, a discoloration is obtained, owing to the previous formation of anilline (commute page 45).

ACRIDINE AND ITS ALLIES.

A cridine and its somer phenanthrene respectively that quinchne bears to apphthalene, and pyradiae to bearcae (compare page 39). The following formulae show their constitution and relationship to anthrecene and phenanthrene:—

$$\begin{array}{lll} C_0H_4 & \left\{ \begin{matrix} CH \\ C_0H_2 \end{matrix} \right\} C_0H_4 & \left\{ \begin{matrix} .C_0H_4 CH \cdot \\ C_0H_1 CH \cdot \end{matrix} \right\} \\ & C_0H_4 CH \cdot \\ \vdots \\ C_0H_4 & \left\{ \begin{matrix} CH \\ I \end{matrix} \right\} C_0H_4 & \left\{ \begin{matrix} .C_0H_4 CH \cdot \\ .C_0H_4 CH \cdot \end{matrix} \right\} \\ & C_0H_4 CH \cdot \end{bmatrix} \\ & C_0H_4 CH \cdot \end{bmatrix}$$

Acridine. C18HON

Acridine has been prepared synthetically by heating concentrated

¹ The base sometimes separates as an oil, which afterwards crystallises.

formic seed on chloroform with diphenylamine and zinc chlorody, and also by various other reactions. Acrdinic is contained in coul-lar, and may be extensed from the fraction boning between 300° and 300°, or from crude commencial antizmeens, by agritaing it with dulute antiphunic and, presuptaing the and liquid with potessium chromate, purifying the scridine chromate by recrystalisation, precipitating the base by animoma, and recrystallization from the twater. The hydrochloride may also be employed for the purification of acrdine

Acridine forms colouriess or brownish-yellow rhombic prisms, of very pungent odour and burning teste. It melts at 107°, sublimes in broad needles at about the same temperature, boils unchanged at \$60°, and distils with the vanour of water.

Acridine is very slightly soluble in cold, but more readily in boiling water, crystallising on cooling in long needles. It is readily soluble in alcohol, ether, benzene, carbon disulphide. &c

Dilute solutions of acridine (and its salts) exhibit a strong blue fluorescence, which is green in more concentrated solutions, and disappears if they are very strong

Certain reactions of acridine solutions with reagents are described on page 126

The most characteristic property of sendine is its intensity irritating effect on the skin and nuceous membrane. Violent sensening and congluing are produced by inhaling the smallest particle of the dust or vapour. The base and its salts attack the tongue even in nunte quantities, and even very dilute solutions cause soute stuging when applied to the tongue or skin.

Acridine has been employed as an insecticide, and compositions containing it have been patented for coating the bottoms of vessels. It is highly probable that the preservative properties of coal-tar creosote oil are partially due to the presence of acridine

Acrdine is a very stable substance. Sulphure and has no action upon it, except at a very high temperature, and caustic potash does not react below 280°. Concentrated nature acid convotrs acidine into nitro-derivatives. Most other oxidizing agents act with difficulty or not at all on acrdine, but by the action

A dordine w best obtained by bestings maximes of one part such of chlorform, diphesylmanse, and muco (preferably) aluminum alboride, with one-half part of ame coulds, for seven or eight hours, under pressure, to 200°-210° C. The points: in bolled with concentrated hydrochloric and, the librard liquid powers dute wester, the liquid again filtered, the executive presentated from the solution by ammonse, and recrystallised from hot water (Fischer and Korner, For., viz. 101). The reactions as is followed:

 $(O_nH_n)_nNH + CHCl_n + ZnO - O_nH_nN, HCl + ZnCl_n + H_nO$

of potassium permanganate it has been converted into quinolinedicarboxylic or aciidinic acid

Acridine is a tertiary amine. It unites with methyl iodide. SALTS OF ACRIDINE

Acrdine is a feeble base. It forms no carbonate, and its salts are more or less decomposed by boiling with a large quantity of water.

Acciding Hydrochlorate, C₁₂H₄N,HCl, forms yallow plates. The solution in water exhibits a blush-green fluorescence, and gives a yellow systalline precipitate of the mescue-ochlorate, (C₁₂H₄N;HCl₁₃HCl₂), an adding mercure chloride With platic chlorate it yields the chloroplatinate, (C₁₂H₂N)₂H₂PtCl₃, in minute, spannigh yallohi, yellow needles.

Avidne Nitité, (Ch.H.N.), HNO, H.O. + 2 aqua, no obtained as a yellow floccible the prespitate of maxing solutions of anithm high-chlouids and sodium nitrite. It forms long, yellow, nilky needles, melting at 1517; somewhat voltate with steam, sightly solution ethni or cold water, more readily in hot water, and very soluble in alcohol

Acridine Sulphito, (C₁₃H₉N)₂₀H₂SO₃, is precapitated in yellowishred or briwnish needles, very slightly soluble in water, on mixing solutions of solume sulphite and acridine hydrochloride, and adding hydrochlorio said.¹

Actidine Pacata, C.311,N.C.3H.(N.C.), This compound is obtained as a canary-yellow precipitate, consisting of minute, yellow, prismate needles, which melt with blackening at 208°. It is almost wholly insoluble in cold, and is partially decomposed by boiling water, it is but slightly dissolved by alcohol or beneue year which boiling. Acidine has been suggested by An a he has he to suggested by An a he has the segment for the determination of pience and, the hydrochloride being used as a precipitant for metallic picates, and a solution of the free base in beneuer for the pience and compounds of hydrochronic being the significant of the precipitant for metallic picates, and a solution of the free base in beneuer for the pience and compounds of hydrochronic.

Hydroacridine. Dihydroacridine
$$C_6H_4\left\{egin{array}{c} CH_2\\ NH \end{array}\right\}C_6H_4$$

This substance is formed (together with a white substance inclubile in alcolol) by the reduction of earchies in alcohole solution by sedum-aunigam. It forms pusses melting at 169°, insoluble in water, slightly soluble in cold alcohol, very soluble in hot alcohol or other. It dissolves in concentrated sulplurace acid, and is precintated unchanged on dilution with water. Augentic and cupies

Before adding send, the liquid contains the compound C₁₂H₀N NaHSO₃, which forms colourless easily soluble pusms

oxides reconvert it into actidine. Hydroacridine is the analogue of piperidine (page 106) and tetrahydroquinoline (page 119).

Phenanthridine
$$\left\{ \begin{array}{ll} C_0H_4 \text{ CH: } \\ C_0H_4 \text{ N} \end{array} \right\} \right\}$$

Phenanthradino is isomoric with scridine, beauing the same relation to phenanthrane that acridine bears to anthracene (P et et and A n k ers mit, Ber, xxii 3339, Jour Sec. Chem Ind., xx 280) It mails at 10^4 and boils about 80° . Henanthritian presents the closest resemblance to acridine, the clufd difference being in its behaviou with reducing agents, for, while acridine yields on reduction a non-base derivative, phenanthridine gives a hydro-base, which crystallises from alcohol in white needles at hydro-base, which crystallises from alcohol in white needles and the same state of the contraction of the new constraints at 235° ; the corresponding compound of phenanthridine at 190° . On adding sodium sulphite to a solution of the hydrochlorade of acridine, a precipitate of reddish-brown needles is produced, while phenanthridine yields no precipitate.

VEGETABLE ALKALOIDS.

The term "alkaloid" was originally applied to the various basic principles existing naturally in plants. As the number of known animal bases increased in number, it became necessary to describe the plant-bases as "vegetable alkaloids" to distinguish them from the alkaloids of animal origin. But with the advance of synthetical chemistry, and the study of coal-tar products, an enormous number of new bases were prepared, and the restriction of the term alkaloid to the natural plant-bases became still more difficult. Discoveries in recent years have clearly established the fact that many of the plant-bases are related to pyridine or quinoline, and several of the alkaloids have been obtained by actual synthesis from pyridine or its derivatives. In other cases, such as canchonine and strychnine, the actual synthesis of the alkaloid has not hitherto been effected, but the relationship of the bases to pyridine and quinoline is not less certain. On the other hand, some of the plant-bases stand in much closer relation to uric acid and the bases found in the animal organism than they do to the other plantbases Thus caffeine and theobromine are undoubtedly uric acid derivatives, while quinine and morphine show no relation to unc acid, being evidently pyridme derivatives

Konigs has proposed to restrict the term "alkaloid" to bases belonging to the second of these classes, and to define alkaloids as "those organic bases found in the plant kingdom which are pyridine derivatives," and it seems probable that this proposal will gradually

be adopted, at least in effect

With the exception of a limited number of volatile alkaloids (eg., nicotine, comine, sparteine), the plant-bases contain oxygen in addition to earbon, hydrogen, and mirogen They are analogues of ammonia, not ammonium bases, that is, they combine with hydrochloric acid and other acids without elimination of water.

The names of the alkaloids are now usually made to terminate in ine, and it is very desnable that this termination should be strictly confined to bodies of a base askura! The termination in is still cumployed for a few of the vegetable alkaloids (e.g., morphis), and by some American writers for certain other alkaloids. The class of bodies known as glue os all es—soms of which are classified an appendix to this chapter, as, from an analytical point of view, they present some similarity to the alkaloids—should receive names having the termination of

The true vegetable alkaloids or plant-bases are very numerous. Many of them are but imperfectly known, while others (e.g., morphine, quinine, streehnine) have been studied very completely.

The alkalords as a class are found in all parts of plants, though in some cases the occurrence of particular salkalods is cursously restricted to certain portions of the plant. Similarly, many of the alkalonis have been met with only in plants of a particular going or family, and in some cases appear to be characteristic of a ungleprocess.²

The vegetable alkaloids are in many cases intensely poisonous (εg) , acoustine, versatime, strychintis), while others, as the alkaloids of coffee, cocoa, and cinchons hark, produce characteristic physicological effects The large majority of them have a bitter taste

With the exception of the non-oxygenated volatile bases, nearly

1 The mass by chemist of the frammation see has caused great confined which it amplicates to designate methanic commencial products jas institution. The size of the size o

3.7. M As a s ch [Phorms Jour. [5], xxx 982, from Jour. Jour. Phon months state that "many off the societies it a similar extensively the class of the state that "many off the societies in the state of the stat

borberine exists in plants of the Remonstration in measurements of the State Radio March 2000 and the Radio March 2000 an

all the vegetable alkaloids are solid at the ordinary temperature. They are in most cases practically fixed, though caffeine and a few others may be sublimed.

Many of the vegetable alkaloads are powerfully sladine are reaction, neutralnee and perfectly, and form well-defined and crystallisable sults. In other cases the base character is only feebly marked, no actuates existing, and even the compounds with the stronger acids being decomposed by mere dilution with

Except the valatile bases, the vegetable alkalonia are, with few oxceptions (e.g., currante, colchume), vary sprangly soluble water, and are consequently prompitated, more or less perfectly, our adding caustie potand or social to the solutions of their sails. In some cases the presputated alkaloni so soluble in excess of the preceptant. The plant-bases can enerty all dissolved by adoption (except rheadme and pseudomorphine), and, as a sule, with great facility. The saits of the alkalonid are usually more adobted in water than the bases themselves, and, as a rule, dissolve also unloaded. This is true of the culphates and other closers of alkalonidal saits, the metallic analogues of which are not soluble in atchied

Certain classes of double salts of the alkaloids (e.g., chloroplatinates, morouro-iodides) are, as a rule, very insoluble in water (compare pages 138, 143)

Solvents immuscible with water differ considerably in their action on alkaloids. The free bases are for the most part soluble, especially in chloroform and amylic alcohol, but in the great majority of cases the alkaloids alie as uncoluble in such mensions as the alkaloids of low base character are decomposed by excess of water, the solutions of these salts often behave with immiscible solvents in the same manner as the fice bases (conpare pages 158, 159).

CLASSIFICATION OF ALRALOIDS

The plant-bases are conveniently stadied in groups, as it is found that the alkaloids of a cestam order or faunty of plants present more or less general resoublance in proposities and composition. Thus the various alkaloids of emchons bank, of opuum, of the accentes, &c., present close analogues among themselves Other alkaloids do not readily admit of being thus grouped, and when of sufficient importance will be described in separate sections

In describing the plant-bases the following general arrangement will be adopted —The general reactions and methods of extracting and purifying alkaloids as a class will first be considered, after your HEARTH.

which the existing knowledge of their constitution will be discussed. The non-expressed volatile bases will then be described. Then will follow sections on the more important supomisable alkalouls, such as the acount and involvate idialouls, and the lakeols of cost. The quium bases will be next considered, and then strychimo and its aline. The cunchous bases will be treated in the next section, which will be followed by one on caffame and its alines. Such of the alkalonds as have not been described under any of the foregoing classes, and which are of sufficient importance, will then be described. In an appeadant to the chapters some of the more important regelable latter principles of non-base character will be shortly described.

GENERAL REACTIONS OF ALKALOIDS.

The plant-bases present more or less general resemblance in their behaviour with certain reagents, and hence their general reactions are classified in the following sections

Reactions of the Alkaloids with Acids.

As bodies of base character, the alkaloids combine with auds to form salts, which in many cases are crystallable and more of see characteristic. They are mostly solube in water and alcohol (including the suil p hates), but insoluble in tallordorm, edges, for Certain of the salts of the alkaloids are safficiently insoluble to allow of the precipitation of the bases for purposes of determinents. Instances of this occur with the picture (bereienen, cumchonne, quinnen), send tatrate (cinchondine), hydroddie (quinnine), chromate (strychnine), hydroferrocyande (strychnine), cauchloride (quinnine, atropine), chloroplatinate (berberne), uniochloride (accontine), and mercuro-oudde (strychnine, neutro, colchierol

Trantios or Alxaloms—In their behaviour with indication of neutrality, the alkaloids, piesent some remarkable differences of behaviour from morgamie bases. The neutral salls of strychning, quinner, morphine, codeine, conine, motine, and other strongly besic alkaloids, are without action on Manss, and these alkaloids can be strated with standard and and littins, just like the morganic bases, except that their high combining weights intensity the effect of the errors of manipulation. Some of the foolbir alkalogids, including nacesine, nacroine, and pepaverine, have no action on intimes, their salts behaving exactly like a corresponding amount of free send.

The salts of the alkaloids with mineral acids are generally

neatral to methyl-organs, which industor can therefore be used to detect and determine any free acid present 1

On nhenolphthalem the great majority of the alkaloids have no action Hence, after neutralising any free acid with the help of methyl-orange, the acid in combination with the alkaloid present can in most cases be ascertained by titiation with standard alkalı and phenolphthalem, and where the combining weight of the alkaloid is known its amount can be calculated from the result of the same titration. The alkaloids to which the process is not applicable are, so far as at present known, atiopine, homatropine, hyoscyamine, hyoscine, and, according to Plugge (Arch Pharm, [3], xxv 45), the volatile alkaloids conine and nicotine. In the cases of brucine, morphuse and thebaine, a red coloration is obtained somewhat before the end of the reaction, but a little experience is stated to surmount this difficulty Moiphine acts as an acid to Pourrier's soluble blue (CLB), probably owing to the presence of the two hydroxyl groups (M. R. Engel, Compt. Rend. on 214)

Lacmond has been used by Van Itallie (Analyst, xiv. 118) for the titration of ceitain alkaloids, including attopine, hyosey-amine and conine, the hydrochlorides of which are stated to be neutral to this indicator

Roselic acid has been employed by E District (Pharm Jow), [3], xvn. 888) for the determination of the alkadeuls in oxtracts of acontic, belladoma, hyoseyamus, contum, and nux vomics, but his results leave the value of the indicator somewhat in doubt

Many of the alkaloids are more or less changed when heated

1 In titrating an alkaloid with methyl-orange, it is larely convoluent to employ an aqueous solution of the base A solution of the alkaloid in proof or rectified spirit is generally suitable, and the indicator is fairly sensitive under such conditions But when the alkaloid is much coloured, as is fiequently the case in the assay of the bases directly extracted from their sources, it becomes difficult or impossible to observe the end of the reaction Under such circumstances, the writer has overcome the difficulty by dissolving the alkaloid in a little other, and placing the solution in a small stoppered cylinder, together with a few centimetics of water, coloured with a drop of methyl-orange solution (1.1000). On then gradually dropping in the etandard acid and agritating tholoughly after each addition, it is easy to observe the end of the reaction, as the colouring matter remains in the upper ethereal stratum, and presents a marked contrast to the red colour of the aqueous hound. By operating in this manner and employing n hydrochloric acid, the author has obtained perfectly satisfactory estimations of acomitine, &c., even when working on as little as 0 030 gramme

with dilute acids, in many cases suffering hydrolysis (e.g., atropine, cocaine, aconitine) or being converted into uncrystallisable isomers (e.g., quinine, cinchonine)

Cont. ' acid, with application of heat, converts certain I' g, morphine, codeine, acontino) into the so-called a po-bases, with loss of the elements of water In other instances, one or more methyl-groups are split off (cocanne, colchemen). For colour-reactions, see page 145.

Concent aded mirric acad oxidises and decomposes the great majority of the alkalods, nitro-derivatives being formed in many cases as intermediate products. In many cases, mirro and yields more or less characteristic colour-reactions with the alkalods (page 146)

Concentrated sulphume and decomposes the great majority of the alkaloids, the change being sometimes accompanied by interesting colour-reactions (page 146) On applying heat, charring frequently ensues. Strychnine survives to some extent a treatment with concentrated sulphume acid at 100°.

Reactions of the Alkaloids with Alkalies.

The fixed alkales, lime, bayte, and ammons, liberate the plant bases from their sails, and as the free bases have, as a rule, but limited solubility in water, they are commonly prequited when the respect is added to their solutions Phebase usually appears as a while, very bulky or floculent prequitate, often exhibiting a crystalline appearance, either at once or on standing The preceptates are often hydrated, and sometimes can only be rendered anhydrous with difficulty

In some cases, the plant-bases when freshly liberated from solutions of their salts by fixed alkales, alkaline earths, or ammonia, are soluble in excess of the precipitant. Thus morphine and codeme dissolve readily in excess of caustic potash or soda, and shighly in ammonia, and morphine is also soluble in lime and bayts water. Quaines, but not other emchane alkalouis, dissolves in excess of ammonia, and strychnine also to a limited extent.

The carbonates of the alkal-metals react somewhat peculiarly with the salits of the alkalonds. Few of the alkalonds form carbonates, so that the preceptates produced by alkali-metal authonates usually consist of the free plant-bases But tho salts of some alkalonds are not preceptated at all by potassum or solume arbonate (e.g., osieune), and others which are thus precipitated are unaffected by bicarbonates (e.g., sirychnine, brucine, attorine, veraturne)

A few of the alkaloids give characteristic colour-reactions when added to fused caustic potash.¹

Saponification of Alkaloids.

Many of the alkaloids, when boiled with a fixed alkalt, baryta, or lime, undergo hydrolyses, with formation of a base of less complex constitution, and the sait of an and usually belonging to the aromatic sense. The change is strictly analogous to the seponification of fats and ethereal saits, and can be effected by boiling with dilute and/s as well as by alkales.

The following equations represent the more important cases of saponification of alkaloids, and show the products of the reaction in each case,—

² Ancolding to W Lean **(Iontebut - steal Chem., xxv 29), onto f? 2 alkalonds examined, only the following gave characteristic colours when fissed with custor potents, 0 of millignames bring med in each cose —Químine, a grassa-green and peculiar oders, quindines, groen, jecenning y allower and finally lowers, necleonaries, or premise with green edge, objecting the bluish-green, enchandines, green, obscipant to propy; cocano, greenish-yellow, funning to blue and durty red on detroper heating.

sulphuric acid

General Precipitants of Alkaloids,

Alkholds as a class give prespirates with a considerable number of resegrate, sepacially compounds of some of the heavy metals. The three prespiration of incline porthaps, a solution of incline of phosphomolykdus and (Somencham's reagent), and a solution of the double notide of mercury and poissaum (Alayar's reagent), but neither these nor any other known segaret will prouptate every alkadord without exception. With the exception of annum, which should be applied in a strictly neithel or faintly alkaline solution, the prespirators for alkadorá should usually be added to a solution of the base slightly aciditated with striphure or acets aced, but in some cases (as in the prescription of cortem piezates) the solution should be strongly aciditated with

Picano Aoro, C.,H.,(NO.), OII. Happe's Braspent. When used as a test for alkaloids, pures and is bost employed in esturated, old, aqueous solution. (I 100). The alkaloidal solution should be rendered distinctly saed with duite sulphure and; except in cases where the alkaloid to be prequitated or sought for so only thrown down in meutral solutions. The precapitated picartee have usually a pale yellow colonu, and are either crystallina or becapes offers since, the forms in many oness being characteristic.

Pieric seid produces no precipitate in solutions (acidulated with sulphuric acid) of aniline, caffeine, conine, morphine, pseudomorphine, solanine, theobromine, or the glucosides, and aconitine, atronine, nicotine, and veratrine are precipitated in concentrated solutions only Atropine and morphine are precipitated from tolerably concentrated neutral solutions Copious precipitates are produced by pieric acid in acidulated solutions of berberine. colchieme, delphinme, emetine, the cinchena alkalords, opium alkaloids (except morphine and pseudomorphine), &c acid is especially suitable for the precipitation of the cinchons alkaloids, and Hager has devised a process of assaying back based on that fact (see Assay of Cinchona bark). Nicotine, brucine and berberine may also be conveniently estimated by piece acid. They should exist as sulphates in moderately acid solution, and the pieric acid be employed as a cold, saturated, aqueous solution, of which 150 c.c. will be necessary to precipitate 1 gramme of the sulphate of a cinchona alkaloid, and twice as much for nicotine sulphate. The following are the limits of dilution at which precipitation occurs, and the characters of the precipitates, according to T. G. Wormley -

ALKALOID				CHARACTER OF PRESIPITATE	Limit of Predictation
Nicotine,			٠,	Amorphous, changing to crystal line tuffs, soluble in nicotine	1 40,000
Conine, .	٠	•		Amorphous, or liquid globules be- coming, crystalline, soluble in comine and agetic acid	1 500
Morphine,				Amorphous	1 500
Codeme, .				Amorphous	1 2,000
Narceine,				Amorphous , soluble in acetic acid	1 5,000
Strychnine,				Amorphous, quickly assuming	1 20,000
Brucine, .				characteristic orystalline forms Amorphous, becoming crystalline	1 10,000
Aconitane,				Amorphous, insoluble in ammonia	1 5,000
Atropine,	٠			Amorphous, changing to very char- acteristic crystalline forms, soluble in weak and, including accide	1 1,000
Veratrine,			٠		1 5,000
Jervine, .		٠		Amor phone	1 1,000
Solanine,				Gelatineus, soluble in oacess of picric acid solution	1 1,000
Gelsemine,				Amorphous	1 500

The alkaloids may be recovered from their pierates by mixing the most precipitate with sodium carbonate, drying the mixture, and extracting with alcohol; or the pierate may be shaken with ammonia and a suitable immiscible solvent

TANNIC ACID precipitates the great majority of the vegetable alkaloids. The precipitates are usually soluble in very weak acids, and in ammonia

The tansates of acontine, brunne, caffeine, colcluene, muphine, physostigmine, and ventrine are dissolved by dilute accite and and tannate of quinne by somewhat stronger and. The tannates of acontine, betwerne, (bruchen), caffeine, unchonine, colchiome, narcotine, papeaverine, thebaine, solamine, strychnine, and ventrine resist more or less perfectly the action of cold dilute hydrochloris and. The tannates of acontine, physostigmine, quinne, solamine, and ventrine are not reclusively below the sulphure acid. Acomitine, physostigmine, and ventrine are completely precupitated by tanne and from solutions strongly activated by sulphure acid, but only partially from elightly aciditated by sulphure acid, but only partially from elightly aciditated by sulphure acid, but only partially from elightly aciditated by sulphure acid, but only partially from elightly aciditated by sulphure acid, but only partially from elightly aciditated by sulphure acid, but only partially from elightly aciditated solutions.

An alkaloid may be recovered from its tannate by mixing; the moist precipitate with recently precipitated lead carbonate or hydroxide, drying the mixture, and boiling it with alcohol

or other suitable solvent, which, on evaporation, will often leave the alkaloid in a characteristic crystalline form

Phosphomolybbio Agid Sonnenschein's Reagent One of the most valuable general tests for alkaloids, and reagent for separating them from foreign matters, consists of a solution of sodium phosphomolybdate in nitric acid. It is prepared by acidulating a warm solution of ordinary sodium phosphate with nitiic acid, and adding an excess of ammonium molybdate solution. The yellow precipitate is separated, washed with water, acidulated with nitric acid, and dissolved in a hot solution of sodium carbonate. The solution is evaporated to dryness and numbed at a low red heat till all ammonium salts are volatilised, the residue moistened with nitric acid, and again ignited. The product, consisting of phospho-molybdate of sodium, is dissolved in ten times its weight of a mixture of one measure of strong nitric acid (sp gr

1.42) with nine measures of water

Sonnenschein's reagent gives yellow, usually amorphous, precipitates with nearly all alkaloids, and as most of the precipitates are very insoluble, a negative reaction with the phosphomolybdic solution affords in many cases a positive ploof of the absence of alkaloids, but, on the other hand, ammonium salts and other nonalkaloidal bodies are also precipitated by Sonnenschein's reagent The phosphomolybdates are decomposed by ammonia, in some cases with production of a white precipitate of the liberated alkaloid. which can usually be dissolved by agitation with a suitable solvent. ieg, chldroform, ether, benzene, amylic alcohol; but when the alkaloid is readily exidisable, treatment of the phosphomolyhdate with ammonia is attended with the blue or green coloration indicative. of reduced molybdic acid. This occurs in the case of aconitine. aniline, atropine, berberine, codeine, colchicine, conine, morphine, meetine, physostigmine, &c Where such reaction occurs the alkaload is best recovered by mixing the moist phosphomolyblate precipitate into a paste with potassium or sodium carbonate, and e4, -1, ig alcohol.

· .. Aoro, Scheibler's Reagent, is used in a similar menner to Sonnenschem's phosphomolybdic solution, and gives very similar reactions with alkalqids. It is prepared by dissolving 100 parts of sodium tungstate and 60 to 80 parts of sodium phosphate in 500 parts of water, and adding nitric acid to acid reaction, or ordinary sodium tungstate may be digested with half its weight of phosphoric acid of 113 specific gravity, and allowed to stand for some days, when phosphotungstic acid will separate in crystals. Scheibler's reagent precipitates 1 · 200,000 solution of strychnine and 1:100,000 solution of quinine The alkaloids

may be recovered from their phosphotungstates in the same manner as from their phosphomolybdates (see above)

Mestatungstic Acid, Silicotungstic Acid (R Godeffroy), and Phosphoantimonic Acid (Schultze) have been proposed as precipitants of alkaloids, but the advantages claimed for them have not led to their general adoption.

BROWINS dissolved to saturation in strong laylve between earth been recommended as a general reagent for alkholois by T G W o'm ley It is probable that hydrochloric acid might be substituted for the hydrobronic acid without datiment to its efficiency. Wormley's Response to others yieldow amorphous procuptates in solutions of many alkaloids, and crystalline precipitates with meconic incidentially strong solutions), actoring, hyosogramme and verstaine, the microscopic appearance of the precipitate being in each case characteristic.

IODINE dissolved in a solution of potassium iodule. Wagner's Reagent, yields reddish or red-brown precipitates with nearly all the alkaloids, even in very dilute solutions. The precipitates are formed more readily in solutions acidulated with sulphune acid, and when applied under these conditions the reagent is in effect iodised hydriodic acid. Excess of the reagent should be avoided The quantity used should not be sufficient to colour the solution vellow Precipitation is so general, and occurs in such dilute solutions, that a negative reaction is conclusive proof of the absence of ordinary alkaloids, though precipitation is not conclusive proof of the mesence of an alkaloid. The precipitates from aqueous solutions are usually amorphous, though codeine, narceine, and strychnine are exceptions. In alcoholic solutions the precipitates are sometimes not formed, or are deposited very slowly, but when produced, they are often of different character from those yielded in aqueous solutions, and in some cases are crystalline. The precipitates are mostly poly-10 dides of the alkaloids, the formulæ in some cases being very complex. Thus with quinine there is first a formation of BHI,I; with more of the reagent, BHI,I, is obtained, while in alcoholic solution, in presence of free sulphuric acid, and

1 C L Bloxam (Chem News, 2iru 1915) has pointed out that contain of the allactoding reclumeterists to holocurrentous when bomme-waters addition by dop to their solutions in dittle hydrochloric and Thus, brucas is stated to yeld a violet colour, and stryphinn the same in louling, narrotine a rose pink, and the same with quanta, changed in the latter case to the characterizing grange goes colour and adding amoness. When the contraction of the co

with an excess of the reagent, the currons 1qdo-sulphate of quanta or her apath 1st, B, 3f, 80, 2MI, I, +3 aq, as produced. Attopine, strychnine, berberine, and piperine are among other alkalonds gramp characteristic compounds with Wagner's reagent. The alkalonds may be recovered from their polypoides by treating the presipatate with sulphurous acid, as sulphite and distinct sulphure acid, or sodium thousulphate, and then adding an alkel and shalong with a suitable immissible solvent. Treatment with sodium thousulphate ("hypochique exces, is a convenient means of purifying the polypoidides from co-prespitated foreign matter. The 1ediced solution is filtered and again treated with Wagner's reagent, when the polypoidide is obtained in a condition of purify.

The strength of Wagner's reagent may vary within wide limits Ordinary decinormal solution of iodine is of suitable strength, or a solution containing 20 grammes of iodine and 50 of polassium iodide per litre may be used

Porassto-lodids or Cadmun, Maimes Reagent, employed in solutions cardulated with sutphieric acid, gives with alkaloids precupitates which are at first anorphous, but which subsequently become crystalline. They are soluble in alcohol, and in excess of the cadmun solution.

Potassio-Iodide of Bisnuts, Dragendorf's Reagent, is best made by mixing 16 measures of the BP solution of citrate of bismuth with 1 of strong hydrochloric and (sp. gr. 116), and adding todide of potassium equal in weight to the hydrochloric acid used (J. C Thresh) The resulting liquid has an orange colour, and when added to solutions of alkaloids, strongly acidulated with sulphume acid, forms orange-red precipitates, which appear to be, in most cases, wholly insoluble in cold water. The following are the limits of delicacy, according to J C Thresh (Phaim Jouin. [3]. x. 641, 809) -Strychnine, 1 in 250,000, quinine, 1 in 200,000. quinidine, 1 in 150,000, cinchonidine, 1 in 125,000, narcotine, 1 m 50,000, brucine and accustine, 1 in 40,000, atropine, 1 in 25,000; morphine and narceme, 1 in 20,000, codeine, 1 in 17,500; apomorphine, 1 in 12,500, berberine, 1 in 6000, caffeine, 1 in 3000. (See also F Mangini, Gazetta, 1882, 155 . Journ Chem Soc, xlu 900)

Porassio-Miraousio Ioduna, Mayor's Reagent, is prepared by distance of the grammes of dry crystallised mercure chloride and 25 grammes of pure potentium todde spearately in water, mixing the solutions so obtained, and diluting the mixture to 1 litrs. The solution thus obtained is grammal, and of convenient strength for general use, though of only one-half the

strength originally proposed by F. F Mayer¹ (Chem. News, vil. 159)

Mayer's Solution precupates the great majority of alkaloids, and in some cases from very ditute obitations. Applied, as it always should be, to solutions tendered distinctly and by hydrochlore or sulphure and, ammount does not interfere, but the colution to be tested must not be more than elightly alcoholic, and must not torain acctic soid. The prequiptates yielded by alkaloids with Mayer's solution are usually yellowish-white in coloni, and outly or floorished. They are more or less soluble in alcohol, other, aceinc acid, includes, and sometimes in an excess of the reagent Certain other organic matters besuited silkohols are also precipitated by Mayer's solution, which therefore loses much of its value whon arrobed to unswingfest solutions.

Mayer's solution is cliently valuable as a means of making an approximate volumetric determination of the alkholog present in a solution; but infortunately the composition of many of the procepitates obtained with twines to a source sextent with the concentration of the solution, the proposition of the and present, and the excess of the reagent

With etychune, the composition of the precupate produced by Maya's collision approximates to BHLHgL, with morphuse it appears to be a variable mixture of R(HI)_{c}(HgL)_{s}, with morphuse it appears to be a variable mixture of R(HI)_{c}(HgL)_{s}, and $R(HI)_{c}(HgL)_{s}$. These formulae refute the estatement made by Mayer, and reproduced by various writers, that the precupitates are of definite composition, containing either 1, 2, on 3 molecules of the base. It has been proved by Ly on shat the precupitates nearly always contain a smaller proportion of mercury (often less than three-fourths) than has been assumed to be present in them. The subject has also been investigated by A. B. Prescott (Chem. Nices, xir 114, 123).

If Mayer's reagent be added till precipitation ceases, there will always be a large excess of the reagent piecent. This excess bears a relation to the dilution of the liquid, and the more dilute the solution, the larger the volume of Mayer's solution requireste

A B Prescott has possible out (Chem. Nines, tw 11s, 123) that the proportions of mecenne and potentium soften used in making Mayar's solution correspond to High-teKI, which implif be supposed to react to form content of the solution point rather to the formain KI, High-teKI, 24CM. I have the reactions of the solution point rather to the formain KI, High-teKI, 24CM. Nevertheless, the proportion post makes the formation of potassims between the solution with more and before the proportion of potassims and potassims bromde, used in the paperties industrial by the formula He(g)+tkHI-tkHI control and potassims bromde, used in the paperties industrial by the formula He(g)+tkHI-tkHI.

effect complete precapitation. Hence, m order to render titution with Mayer's solution of any value, it is securated that the solution operated on shall be nearly of uniform strength, and that the research be added in exactly the same manner. It is futther desirable, whenever possible, to make an experiment, sade by side with the alkalonida solution, with a known weight of the same likely and in a state of purity, so as to avoid all assumption as to the behaviour of the volumetic solution with the alkalonid analonism.

The following is the usual method of performing the thrinton of an alkalod with Mayer's solution. —The solution, which should be distinctly accidated, and contain, as a rule, 0.5 per cent of the alkalod, is irrested with Mayer's solution as long as a distinct precipitate is produced. As there is no definite out-reaction, and no astafactory indicator has been as yet devesel, it is necessary to filter a portion of the solution to ascertain if the precipitation is complete. A munute filter, about half an inch in diameter, supported on a ring of pishumu-wirs, may be used. A drop of two of the filtered liquid 2 is pieced on black giasa; or on ordinary glass on black paper, and a drop of the volumetrie solution added from the burstet, when the fantest turbulity will be readily preceived. Before the end of the timuton, all the trial-filters and test-drops are returned to the solution constaining the man quantity of the precipitation.

The end of the reaction is the point at which the Mayer's solution ceases to produce a precipitate, and it is worthy of notice that, before this point is reached, a condition of equilibrium is etianted, in which the solution is liable to be precipitated by the addition of either alkaloidal solution or the mercury research.

A B Ly on a has investigated the behaviour of vanious alkaloids with Mayer's solution, noting the effect of concentration said the volume of the reagent required to prequitate completely a definite weight of alkalod, in adultion, the volume required produce an apparent excess of the mercury reagent (so that the liquid would give a preceptiate with more of the allacloid action), and also the actual excess of Mayer's solution used, as estimated from the quantity of mercury present in the solution

Lyon's results are given in the following table, reproduced from his Manual of Pharmaceutical Assaying The mercurial solution was an normal, and 0 1 gramme of alkaloid was employed in each case—

¹ F F May s; proposed to sesertum the excess of the teagent by tritating back with standard mirate of silves solution, without filterings using potassium chromate as an indicator. As pointed out by Lupinski, the suggestion ignores the secuminishton of chlorides and includes in the solution, as size the fact that some of the procupitates reach the slowly with intrate of silver.

² A convenient form of filter-tube for the purpose has been described by F. C. J. Bird (*Pharm Jour.*, [3], xvii 826)

Alkaloid.	Solu	tion	Volum	in c. c.	agent	Weight of Alkaloid	Weight of
	Condition.	Strength	For apparent excess	For com- pleto pre- cipitation, execis		prozipitated by 1 c c of Resgent.	Precipitate
Aconitine, Atropine, Berberine,		1 200 1 200 1 400 1 400 1 200 1 400 1 600	7 0 6 0 6 9	7 1 13 1 14 0 16 0 8 8 3 9	2 0 8 0 8 5 8 0	0141 0/77 0/72 00/72 00/07 02/08 02/77	180- 190 216- 220 192- 200
prucine,	Nearly neatral	1 200	1 1	4 0 8 0	17	0218 0125	200-215
,,	Nearly neutral Acad	1 400		88		-0114	
Ciuchonidine,	Nearly neutral	I 900		98 92		0102 0109	
Cinchonine,		1 100 1 200 1 200 1 100	12 4 12 4	18 8 18 5 16 0 12 8	1-0 07 2-6 0-8	0078 0074 0001	880- 875
" "	Neutral Acid Neutral Acid	1 100 1 200 1 200 1 100 1 400	7 9 8 0 8 0	14 0 10 8 14 2 12 4	12	0078 0072 0008 9071	883 845
Cocaine,	1	1 200 1 400 1 600	10 0	14-18 12 8 14 4	40	007 to 10088 0078 0009	246
Colchicine,		1 200 1 400 1 000	8·2 4·2 5.0	16 0 9 2 11 4	52	000S 0109 0088	160
Emetine, .		1 800 1 200 1 400 1 600	4 0 8 9 8 8	14 6 0 4 10 2	01	0090 0097 0106 0098	258
Hydristine,	. 1	1 200 1 400 1 200	5 8 9 5	10 9 10 4 12 0 7 4	08	0004 0006 0084	185- 200
Iyosoyamine,		1 400 1 600 1 200 1 200		8 0 8 4 8 5		0185 0125 0119 0110	200- 210
locerpine,		1 400	7-0	4 01 8 9 16 8	00	0128 0110	220- 250 100- 210
uinine,	Neutial Aud	1 200 1 200 1 200	11 6	20 0	1	0060 0060 0001	240- 850
taychnino,		1 400	12 8 12 2	18 0 16 8 20 0	- 1		810- 885
"	Acid Acid	1 200 1 400 1 400	11 6	11 0 12 0 12 2	0.8	0001	275
		1 000	11 2		08	0082 0087	:

From a study of this table by Lyons, it appears that while a notable excess of the reagont is generally needed to effect complete precipitation, the weight of the precipitate is in many cases considerably below the amount inducated by theory. Better respect are obtainable by, allowing the liquid with the suspended precipitate to stand for some time. Lyons states that, under these circumstances, the atropine presuptate becomes dense

and crystalline, and in part adheres to the beaker, in which it can be washed by decantation, dired, and weighed, the amount thus found falling little short of the theoretical weight of 0.245 gramme for 0.100 of alkaloid

The following data showing the behaviour of alkaloids with Mayor's solution are tabulated from the descriptions of Dragendorff (Plant-Analysis and Analyse Chamique de quelques Drogues Actives) —

Alkalold.	Dilution of Solu- tion	Multi- grams of Alka- loid Ppted by 1 c c	Correction for Solu- bility Mgrain for 10 a.o Filtrate.	Observes	Conditions of Preceptiation
Acontino, Proudacinitios, Atropue, Broscyamine, Broscyamine, Broscyamine, Strichnine, Pastrine, Sobathline, Sobathline, Britanine, Britanine, Britanine, Schathline, Schathlin	1 200 1 330 1 200 1 200	17 4- 19 4- 19 4- 19 4- 19 4- 19 4- 19 6- 20 6-	05	Dragendorff " " " " " " " " " " " " " " " " " "	Ample time sequired for measurements. Faintly send only KCI present and Label and Lab

H or s h (Phasus, Record, 1886, page 209) has proposed an improved method of operating with Mayer's solution, which ellows time for the paceiptate to fully form. A number of equal portions of this solution to be tested ane treated with volumes of the mercurial solution, regularly increasing by 0 1 c, and allowed to stand eight or ten hours. That-positions of each nuxture are then removed and bested with two drops of Mayer's solution, when a putticular nuxture will be found to have the mecurial solution in slight excess, while in the previous muxture it is deficient. Obviously, the true amount lies between the two, and it is easy to ascortan the exact volume required.

Strychnine and quinine are among the alkaloids yielding the most insoluble precipitates with Mayer's solution. With atropine

and morphine the reaction is far less delicate, and caffeine and theobromine are not precipitated at all

MERCURIC CHLORIDE, HgCl₂, gives, with certain alkaloids, precipitates of which the crystalline form of melting-point is characteristic. As a rule, the precipitates have the constitution B₂H₂H₂Cl₄ and are less insoluble than those produced by Mayor's respect

Acuse Chicanna, Ancil, grees yellow preceptates of alkaloidal auto-chicardaes or chicarus ates with hydrochlore and solutions of many of the alkaloids The double sales preceptate are often very modelule They usually contain B, HGLAudl, or BHAudl, though this formula is not without exception. Auric chlorido has the advantage that ammonium sales and the simpler anides are not precipitated by it, but the precipitates are unstable, the yellow colour in many cases rapidly changing to reliably brown, while the supernatant liquid occasionally sequires an intense red colour

PLATING CHLORIDE, PtCl., is a usoful reagent for many alkaloids, with the hydrochlorides of which it combines to form chloroplatinates or platinochlorides. In some instances, these double salts have the formula BH, PtCla, and m other cases they contain BoH.PtClm while in a few instances more complex formulæ have been attributed to them. It is sometimes stated that the alkaloids containing No in the molecule form chloroplatinates of the first formula, while in the case of bases having only one atom of nitrogen the platinum salts contain two atoms of alkaloid, in other words, that the iatio of N Pt is constantly as 2 1. This, however, is far from being the case, for alstonine, golsamine, aspidospermine, paytine, strychnine, pilocarpine, and numerous other bases containing No agree with the opium bases, berbeijne, cevadine, atropine, and others containing N in forming platinum salts of the formula B2H2PtCla. In addition, many of the cinchona-bases form platinum salts of both series

The chlocoplatmates of the alkaloids vary in colour from pale yellow, though orange and rod, to howmash red. They are mostly spannegly soluble in water, and hence are usually formed as preceptates on adding platma chloride to a solution of the alkaloid acidulated with hydrochloric scal. The similar behaviour of potash and ammonia drammabes the value of the test. Xanthame, orferine, solutione and pelleteriem are among the alkaloid not precipitated. Of the rest, the chlocoplatmates of quinne, cinchonion, morphine and sulvehime are among those dissolved by hydrochloric acid. The melting-points of the alkaloidal chloroplatmates are often characteristic

Potassium Permanganath, KMnO4, produces characteristic

reactions with certain of the alkaloids. Beckurts and List have examined the behaviour of a number of them, by adding a domnormal solution of the respent, drop by drop, to a cold startisted aguscous solution of the bydrochlorate of the bare limited the restriction of the permangansky with separation of brown manganese coxide, was observed with the hydrochlorates of brown manganese coxide, was observed with the hydrochlorates of quintie, einchonlande, einchonnine, einchonnine, broncie, versatine, colchiens, contine, contine, physostigmine, todeun, and thebaine The solutions of stropine, hyosyamine, pilocarpine, berberne, piperne, and strychnine wore coloured red, the resgent bang only gradually reduced

With morphine hydrochlonds the permanganate produced a winte crystalline precipitate of oxydimorphine, which, when fillored off and drad, could be recognised by its characteristic reactions. Apomorphine hydrochlonds immediately reduced the reagent, with moduction of an intense erren colour

On adding a few drops of a decinormal solution of potassium permanganate to a concentrated solution of naresume hydrochlorida a reddish precupitate is numediately formed, which is very stable in the sold and in the absence of an excess of the reagont, but is decomposed on heating or by addition of more permanganate Solutions of papaverine hydrochlorida, and of narootine if ultitude with hydrochlorine soul, at first behave similarly, but the preciptives are much less stable than narosine permanganate, and soon discolour and decompose with separation of from manganess exide,

F. Gersel (Phām Zeit, 1886, p. 132) has pointed out that occause gives a comparatively stable permanganate, which forms a purple-violet precipitate of characteristic microscopic appearance. The precipitate forms only slowly in dilute solutions, and undercose gradual decomposition.

Colour-Reactions of Alkaloids.

Many of the alkalouds give brulhant, and m some cases characteristic, colorations when treated with appropriate reagents. When possible, the reaction should be compared with that yielded by the pure alkaloul treated side by side with the sample. The reagents which have been proposed as colour-tosts for alkalouds are very numerous, and have not always been chosen or applied with discretion, nor with a due regard to purity. The colour-reactions may be classified as:—(1) Those produced by dehydrating agents, such as strong sulphurne and, phosphorosicit, and sure chloride, '(2) those given by oxidinging agents!

¹ In using one chloride, Czu mpelitz directs that the substance to be examined should be first exceptilly dired, moistened with a solution of 1 gramma

not of themselves yielding colours, such as intre eard, chlorine, bromine, and bleaching powder, or stiphure and and oxidising agents, such as potassium chlorate, perchlorate, and permanganate; (3) those given by oxidising agents which themselves yield a coloured product by reduction, such as notice and reagents containing chromic, molybdie, tungste, and vanadic acids, (4) and colorations produced by certain special tengents, such as ferrice chloride, hydrochloric scale, sulphure acid and sugar; &c.

As a rule, the best method of observing the colour-neactors of an alkalond is to anply a drop of the reagent by meens of a pipette or glass rod to a minute fragment of the solid alkalond, placed on a poscelam plate or in a flat poscelan dish Ari, alkalondal readule obtained by the ovaporation in a porrelami; capsule of an alcoholo, ethersal, otheroforme or other solutions may be vary conveniently omployed for observing colour-reactions;

Fusion Caustro Forase gives a few interesting colour-reactions with alkaloids (see foot-note, page 133)

Coxermination of the Actor gives colour-reactions with a few it could be a few a few and vestignine, sabatiline, vestirine, and vestatordine, and a yallow with thebame On addition of chlorine-water after hydrochigne, and, berberne gives a sed colour. Nicotion spitch an amorphous hydrochloride and comins a crystalline salt, on evaporating the solution in hydrochloric acid

CONCENTRATES SULFERING ACID gives colour-reactions with a number of alkalonds, the coloration varying with the degree of heat applied The following reactions have been observed when the card is dropped on to the soil alkalond, without applying heat —No colora, or a faint straw that only, is yielded by pure acounting, actroping, edifferent, cholonding, enchounding, cocaine, codeme, hyoseme, hyoseyamine, gelseanine, morphine (juriple to hown on warming), nechine, pulcostraine, quinne, quinnel, quinnel, are given by colorations, groscopine, jervine, and by many other alkaloids in presence of impunities Reddist colorations are given by colorations, groscopine, jervine, and by many other alkaloids in presence of impunities Reddist colours are produced either immediately or gaidually, with impure acontine,

maissi zun, chiorndo un 30 ce of water, and duvel agaun If thus treated, strychune takes a scalet colour, thehame a yilow, anacene in diver green, delphinnien red-brown, batheme a yillow ventaine a red, quinna e pale yillow, digitalin a manoon, saluan a rudel-red, santonin a vioqui-thus, and cubelun a purple. The puesmes of binnen prevents the coloration of skychnne, the tingo produced being a duity yellow (Giornale Farm Chem , Jour Chem Soc., Xii 1840)

¹ Information respecting this test will be found under "morphine."
VOL III PART II. K

apomorphune, brucune (mle rose), oceano (unpure), conna (quale red), gaisemme (unpure), mecondine, narveue (changung to black), narcotine (yallowsis-red, changung to vtolet and blue), physostagmune, hickenduce, subdullune, abutatune, solame, taxue, thebaue, veratune, and verestodine. Blusis colombions are yielded by errjptopine, currente (after at tune), and paparente Greenste holder are given by beberne, borberne, enseine (trownash to green), piperine, pseudomorphine, and sonestimes by phroadune.

Some characteristic changes of colour can be obtained by gradually warming the capsule in which the test is being mode, by placing it over a small beaker of boiling water. The ultimate result is usually browning and chairing of the alkaloid, but the intermedual caections are often of value.

Many substances besides alkaloids give more cless bulliant colour-meastons with stong subplume and Thus sed colorations (often of a bulliant hue) are obtained with amygdin columbin, cubbin, clatern, besperdin, pholoran, populu, salten, sarsayaullin, senagin, smilacin, syringin, and many varieties of tannin

In applying sulphuise acid as a colour-test for alkaloids, it must be remembered that the presence of a very minute quantity of nitiic acid, often present as an impurity, greatly modifies the colorations produced by many of the alkaloids. Thus, if the treatment with sulphuric acid (without applying heat) be followed by the addition of a very minute quantity of nitrie acid (at the end of a glass rod drawn out to a point), or a minute fragment of solid potassium nitiate, the following reactions will be obtained 1 No coloss with atropine, caffeine, cinchonidine, cinchonine, nicotine, pilocarpine, quinidine, quinine, staphisagrine, strychnine, or theobromine, red coloration with brucine, curarine, narcotine (reddish violet or blood-red), physostigmine, sabadilline, thebame, and veratrine (gradual change to cherry-red). Special and peculiar changes of colour are produced by morphine, codeme. and colchieme, and are described in the respective sections on these alkaloids

STRONG NITRIO ACID, of 140 to 142 specific gravity, gives more or less characteristic colour-reactions with a number of alkaloids. A drop of the acid should be applied by means of a glass

¹ Er d'mann apples thus test by mixing 6 drops of nutre aud of 1 as species gavathy with 100 o. o. of water, and skining 10 drops of the dutus sold so obtained to 20 grammes of subsure call From 8 to 10 drops of the solution so prepared, or Fortunatia Report, is added in 1 or 2 million grammes of the solution to be tested, and the colour observed after 20 to 30 minutes.

rod to a minute fragment of the alkaloid, or to a residue left on evaporating a solution on white porcelain No coloration is yielded by aconitine (when pure), atropine, caffeine, cinchonidine, cinchonine, conine, gelsemine (impure, gieonish), quinidine, quinine, strychnine, or theobromine. Yellowish colours are obtained with impure aconitine (colour varies from vellow to red and brown). codeine (orange-vollow), morphine (vellow to red), narceme, narcotine, papaverino (orange), piperine (orange), rhosadine, sabadillina (vollow), thebaine, and veratrine Red shades are produced by impure aconitine (colour varies from vellow to rod and brown). anomorphine, beharine (red to red-brown), borberine (red-brown), brucine (blood-red), papaverno (orange-red), pscudomorphine (orange-red), and physostigmine (gradually) Gelsemine yields a deep bluish oreen residue on evaporation. Blue colours are said to be given by colchicing and solanine, and by the glucosides igustrin and syringin

SULPHOMOLYBDIC ACID, Frohile's Reagent, affords one of the most useful of the oxidation-tests for alkaloids, but it must be borne in mind that the colours produced are in great measure those of the lower oxides of molybdonum, and that various other bodies besides alkaloids readily reduce molybdic acid with formation of these coloured oxides. The reacent itself, if strongly heated, acquires a blue coloration from reduction of the molybdic acid Frohde's reacent is prepared by dissolving 5 milligrammes of molybdic acid or molybdato of ammonium in each 1 cc of stiong sulphuric acid No colour is produced with atropine, caffeine, cinchonidine, cinchonine, conine, delphinine, hyoscine, hyoscyamine, meeting, strychning, or theobioning, Yellowish colorations are given by aconitine, colchicine, and piperine. Reddish shades of colour are produced by brucine, emetine (red, changing to green). narceine (red, changing to blue), sabadilline (reddish violot), solanine, thebaine (orange), and veratrine (gradual production of a cherry-red colon) Blush colours are given by codeing (gradual moduction of deep blue), morphine (violet-blue, then dirty green, changing to deep blue), narceme (yellowish brown, changing to red and blue), staphisagrine, (violet-brown) Greenish colorations are produced by apomorphine (green to violet), beberine (brown-green), berberne (brown-green), emetine (red, changing to green, and turned blue by hydrochloric acid), quinine (pale green), and quinido (palo green)

Of non-alkaloidal bodies, colocynthin gives slowly a cherry-red colour; olaterin, a yellow, phloridam, giadually, bline, populin, violet; salicin, violet to cherry-red, and syringin, a blood-red to violet-red coloration.

Sulphovanadio Acid has been recommended by F Kundrát (Chem Zest, xiii 265, Jour Soc Chem Ind., viii 421) as a colour-test for alkaloids The reagent is prepared by dissolving 0.1 gramme of ammonium vanadate in 10 cc of strong sulphuic acid. It is stated to give the following reactions, many of which are due to the production of the coloured lower oxides of vanadium, and hence are likely to vary with the proportions of the reagent and alkaloid employed No coloration is produced by caffeine or nicotine Brown by acomitine (light coffee-brown), codoine (greenish brown, becoming darker), morphine, naiceine (changing to diriv blush violet, then gradually reddish brown), piperine (intense reddish brown to black), kairine (dirty pink, quickly changing to duty light brown and brownish green), solanine (coffee-brown, changing at the edge to purple and in the centre to dirty green, and very gradually becoming an intense violet jelly) Red colorations are given by stropine (changing to yellowish red and yellow), brucine (intense blood-red, gradually fading), narcotine (blood-red or purple), and veratrine (biownish red, changing to reddish violet). Yellowish or orange colours are produced by cinchonine (changing to green), cocame (orange, froths on dissolving), physostigmine (greenish yellow, then purple, finally yellow-brown), and quinine (changing to bluish green and greenish brown) Green colorations are produced by colchicine (changing to greenish brown and coffeebrown), conine (intense green, changing to brown), and quinidine (faint bluish green) Blue reactions are produced by antipyline (greenish blue, gradually becoming bluer), and apomorphine (dark violet blue, rapidly changing through duty green to reddish and hght brown) Violet colorations are given by papaverine (gradually changing to bluish green and orange-green), and strychnine (bluish violet, changing to reddish violet, purple, and brilliant red)

Of colorations with non-basic principles the following have been recorded.—Antifebrin, purple, rapidly changing to hown; given tailin, intense brown, with reddish shade, and salvejtic acid, brownish green Preretoxm and santonin give no coloration with sulphovanulos acid

FRENCO CHICARINE gives a few characteristic colorations, the most important bung the blue reaction with morphine and the blood-red with antipyrum (page 35) A freshly-made unixture of four ic bilor de and potaesium (perrugunade (free from farrocyanide), both in aqueous solution, has a yellowish buvon colour, yellow presence of certain alkaloids it is immediately coloured blue (or green) owing to the production of Plussania blue. This reaction was at first regarded as characteristic of the pto tomaines or conductor to bease, but it is produced by any readily oxidishle

alkaloid, and hence is given immediately by morphine, acontine, physostigmine, &c, and after a short time by hyoscyamine, emetine, colchicine, nicotine, and many of the tar-bases

OUDLATION-OBLOTH-READTIONS are also produced by resignits abruing a none powerful oxidising action than the foregoing. Thus strong sulphune and may be employed in conjunction with potassium initials, chlorate, perchiorate, permanganate, biolinemate, and ferineyandle, on with melhilic perceives, such as those of manganess (MrO₂), lead (PbO₂), rathenium (RoO₂), unanium (U₁O₃), and ceium (Ce₂O₃). The nost important of the colour-reactions obtained with such reagents are those yielded by strychnine, curing calesium and anihus, which are fully ideembed elsewhere.

Physiological Tests for Alkaloids.

A large number of the natural alkaloids, if not an actual majority, have well-marked poissonous characters The symptoms produced are of the varied description, ranging from the narcotism of morphine to the paralysis of contine and the tetanus of strebnine

In making experiments on animals it is often advantageous to administer the poison by hypodermic injection of a solution of alkaloid in water, or weak spirit acidulated with acetic acid. Such a plan obviates the loss of the poison by vomiting, which sometimes eliminates the greater part of the poison from the system On the other hand, the subcutaneous injection of small animals as onen to certain obvious objections, and in many cases internal administration may be advantageously substituted for it, especially if the animal employed be a rabbit or guines-pig, and hence not hable to vomit. In many instances, such animals are hopelessly large, and mice, small birds, or frogs must be employed Wynter Blyth has used blowfiles with success in some cases, and occasionally fish are of service. When the poison is to be given internally, the extract or very strong solution should be made up into one or more small pulls with oatmeal, which the animal is either induced to eat or forced to swallow In the case of hunets and other small birds, a drop of the liquid to be tested should be introduced into the open beak by means of a pipette or feather

In some cases, physiological tests may be advantageously made on human subjects. Besides observing the hitter taste possessed by most alkalouds, the tingling sensation produced on the tongue by acountine and cocaine can be thus detected

A marked physiological characteristic of many of the alkaloids, sufficiently striking in some cases to serve as actual evidence of their presence, is their effect on the pupil of the eye. The test is generally made by placing a drop of the alkaloidal solution to be examined, as nearly neutral as possible, on the eye of a rabbit, dog or cas, when, in a time varying from a few minutes to about half an hour, a marked contraction or dilation of the pupil will be observed.

A. The pupil is dilated by :--

- 1 Atropine and belladoma, hyoscyamine and hyoscine, and preparations of henbane and stamonium, solanine, and extracts from solannocous plants generally.
- 2 Cocaine, and preparations of coca
- 3. Conine, and preparations of hemlock and other umbelhierous plants
- 4 Cytisine, and preparations of laburnum.
- 5. Digitalin, and preparations of foxglove.
- 6. Gelsemine, and preparations of gelsemium (yellow resamine)
- 7 Sparteine, and preparations of broom
- 8. Veratrine, jervine, and preparations of hellebore
- 9. Hydrocyanic acid and cyanides.

Mydracus, or diskton of the pupil, is so striking a characteristic of atropine and the isomeric and associated bases that these nor often grouped together as the "mydrastic alkaloida." The mydrassi is only observed in the eye to which the alkaloid is applied

- B The pupil as contracted by .-
 - Morphine, and other opium alkaloids and preparations of opium.
 - A conitine, and preparations of aconite and other members of the Ranunculacea
 - 3 Physostigmine, and preparations of the Calabar bean
 - 4 Strychnine, brucine, and preparations of nux vomice

A similate affect on the pupil is produced by the poisons when taken internally or hypodermically in sufficient quantities. Sometimes, as in the case of morphine and preparations of opium, the pupils are continuted during the early stages of the poisoning, but dilated subsequently, especially after death. Nicotine and preparations of tobacco in some cases cause contraction, and in others dilaton, of the pupil. In poisoning with a contribution, and contraction and dilaton of the pupil is constrained observa-

ISOLATION AND PURIFICATION OF ALKA-LOIDS.

The vegetable alkaloids are found in all parts of plants, and in many cases constitute their chameteristic active principles. It must not be assumed that the active principle is necessarily of an alkaloidal character, though plants and plant-hyordicat, which act primarily on the nervous system, producing tetenus, paralysis, or narcoss (eg, nux vonines, acontie, opium), over their activity, as a ratio, to the presence of an alkaloid. On the other hand, in plants which act primarily on the nususular system <math>(eg, digitale), the active princip is usually of a non-alkaloidal chanacter. Whee the action of the plant is emetic, exhautic, or pruchy astrugent, but this statement has some marked exceptions, for processanlar, a stypical emetic, owes its activity to the alkaloid on in in e

An akkalond never exists n a plant in a free state. It is most requirely revent as a sale, do then an and ash, of some organic and, especially malic acid or one of the varieties of tannic acid. In some instances the acid with which the akkalond is unted is peculiar to the plant in question, as, for instance, meconic acid in new popular, quinic acid in cutchions balk, and igs auric acid in new commerce. In other cases the akkalond is combined with an inorganic acid, as is the case, in part at least, with the morphine in optime. The natural forms of combination of the akkalonds are almost invariably reachly soluble both in a water and in alcohol, but insoluble in effer

The general action of solvents on the leading constituents of plate is the seen from the following table, which will also serve to indicate the nature of the bodies lakely to be co-extracted with the alkaloid when the respective solvents are employed —

	Water	Alcohol	Ether	
Alkaloudal saits, Other saits of inorganio soids,	Soluble Mostly soluble	Soluble Mostly maniable Soluble	luzolubie Insolubie	
Other salts and organic acids, Free organic acids, Tshnins and colouring	Soluble Soluble, Soluble	Soluble Soluble	Mostly macluble Mostly insoluble Variable	
matters, Sugars, Gums and pectous bodies, Albuminoids, &c , Starch.	Soluble Soluble Soluble Soluble Soluble in hot	Soluble Mostly insoluble. Insoluble Insoluble	Insoluble Insoluble Insoluble, Insoluble	
Cellulose, Realns, Fixed oils,	water Insoluble Insoluble Insoluble	Insoluble Soluble Sparingly soluble	Insoluble. Variable Soluble	
Essential olls, Chlorophyll,	Insoluble Insoluble	Soluble Soluble	Soluble Soluble	

Alcohol as the solvent best adapted for the extraction of alkiloids from plants, which should, of course, be reduced to a suitable condition. The treatment may with advantage be repeated several times, the reaches being well pressed between each extraostion, which is preferably effected by a percelutor, or some equivalent arrangement. In the final extraction, the addition of a hittle sulphinic or tertaine and is often an advantage, but the amount of and used should be very limited, and its employment is vetoed in the case of readily clanification alkiloids. Hot water may be substituted for sloohol in some cases. When alcohol has been used for the extraction, it should be removed partially or wholly by goalle evaporation before proceeding to the next stope of the treatment.

The method to be adopted for the solation of the alkalond from the infission or tructure obtained depends much on its nature, and the object of the experiment Extraction by immercible solvents permits the detection of small quantises of alkalonds, which defy methods based on prescriptation, and hence this principle is very valuable in toxicologous investigations; but, on the other hand, the alkalonds so extracted are usually less pure than when isolated by other means

Where it is intended to attempt the separation of the alkaloid by conversion into an insoluble or nearly insoluble compound, a variety of precipitants are available, each one of which has special advantages in particular cases. But before resorting to these general precipitents, it is desirable, and in many cases absolutely necessary, to remove from the liquid as much as possible of the inert organic matters. The best resgent for this purpose is lead acotate, which should be added gradually to the previously neutralised liquid, as long as a precipitate continues to be produced. avoiding the use of any considerable excess of the reagent. The precipitate having been filtered off, the filtrate should be treated with basic acetate of lead, which in many cases will produce a further precipitate, to be removed by the filter as before. On adding ammonia to the filtrate, a third precipitate will frequently be produced, but it must be remembered that cinchonine and other sparingly soluble alkahes are hable to be thrown down at this stage.1 (On this account it is undesirable to add basic acetate of lead and ammonia at once, and filter off the joint precipitate.)

¹ The threefold treatment with neutral lead acetate, base lead acetate, and ammonia in presence of lead acetate causes the precipitation of taninas, most vegetable acidis (e.g., maile, to tarin, oxide); albuminoside, starches, and guns, many glacosides, sugars, and devirin; and the majority of colouring matters

The huuid, which should smell distinctly of ammonia, is next evaporated at a goatic heat tull the olour of ammonia has disappeared, when the excess of lead is precipitated by a stream of sulphureiged hydrogen or the addition of a moderate excess of ditties sulphure acid. Of these plans, the first is much to be preferred. The lead sulphule often carries down with it a notable quantity of colouring matter, otherwise difficult to remove, and the excess of sulphuretted hydrogen is easily got ind of by concentrating the filtrets at a gentle heat. When sulphure acid has been employed to precipitate the lead, the filtrate should be carefully neutralized before attempting to further concentrate the head, otherwise the alkaloid may suffer partial or complete decomposition

The alkaloudal solution, having been purified by the foregoinfor treatment, may be treated with one of the general regently as alkalous, the choice of which will necessarily depend on the nature of the base supposed to be present. Where this unknown, preliminary tests with various precipitants should be made on small adopted fractions of the solution. Although other reagents may be preferable in particular cases, the choice will generally he between one of the following prepulpitants:—

- A fixed alkals, carbonate of alkals-metal, Isms, or ammonia; suitable for precipitating morphine, the cinchona alkaloids, the acouste bases, &c.
- Picric acid (page 134), very suitable for precipitating the cinchone bases, emetine, berberine, and verature
- 3 Tannic acid (page 135)
- 4 Phosphotungsite or phosphomolybdic acid (page 136), available for the great majority of alkaloids, and especially for strychime
- 5 Iodised rodide of potassium (page 137), which produces very insoluble precipitates with the great majority of alkaloids
- 6 Mayer's solution (potassio-iodide of mercury) (page 139), valuable for precipitating emetine and the option bases.

With the exception of tanine acid, which should be applied to the neutral or even family alkalms solution of the alkaloid, the reagent should be added to the acidulated solution, subplume acid being the most suitable acid to bring the highest lot bits proper condition. In most cases preceptation is tolerably reach tit is desirable, as a precention, to wait 24 hours before proceeding with the fiftration. Thus is especially necessary perhaps in the case of precipitants 1 and 2. The alkaloid may be

recovered from the precipitate in the manner described on page 135 et seq

As a rule, the salts of the alkalouds are not soluble in immuseable solvents, and honce when the accidulated solution of an alkaloud is existed with chloroform, either, petroleum sparal, benzene, or amylic bloobel, the solvent does not remove the base from the aqueous liquid. This behaviour breadly distinguables a lk aloud is from glucosides, but, owing chiefy to their weak basic character and the instability of their sells, cafferen, coldenies, delphinuse, narcotine, papewerne, thebane, and theolucime are partially or wholly removed from their acidulated solutions on aguitain with chloroform, while amylic alcohol is stated to extract berbeine and ventures in addition to the above bases

EXTRACTION BY IMMISCIBLE SOLVENTS

The behaviour of the alkaloids, when their soid and alkaloid solutions are agitated with immiscible solvents, is of the highest practical value for their isolation and identification.

The numesoble solvents used for the extraction of alkaloxia, &c., ahould be tree from any tance of fixed or difficulty volation organic matter. This is best ensured by shaking the solvent with water alightly sciulated with subplume acid, separating the squeous layed, and relatifiling the numesoble solvent at a moderate tentral content of the solvent at a moderate tentral content with the solvent and indeed may be advantageously kept in contact with family alkaline water. The agriculture with vater is essential in the case of solvents liable to caning abook (e.g. either, chloroform, publication), the presence of which might seriously modify their factors.

In using mnuscable solvents, it must be borne in mind that extraction is nover theoretically perfect with a single treatment. The dissolved body is distributed between the two solvents in proportions which are probably dependent on the islative solitility of the substance in the two media, and the relative quantities of the two media employed. Thus, it may be supposed that if a substance be 99 times more soluble in chloroform than in water, and its aqueous solution be shaken with an equal

¹ The principle appears to have been first adopted by Otto in 1556, who employed other in his modification of 8 tars 'process for the detection of possenous sikelands. In 1856, Bed gers and Otto wood amployed the method with chlorofers, and in 1811 Usia and Erd ma na recommended the use of surplus should in 1807, Dragon for if pubblished has well—the state of the process of the state of the process of t

measure of chlomform, 99 per cent of the whole substance will pass into the chloroform. On squareting this layer and soon amtatang the squeous bound with an equal quantity of chloroform, 99 per cent of the remaining substance will be dissolved. thus making the exhaustion practically complete. In the case of other and ample alcohol the solubility of the solvent stealf m the acrocons liquid is also an important consideration, for as ether is soluble in about ten times its measure of water on agitating together sonal measures of other and an agreeous bound it may be assumed that one-tenth of the other will be dissolved. and will remain in the aqueous hand together with its ones tenth share of the alkaloid or other substance to be extracted. On separating the othercal layer, and again shaking the someons hourd with an coual measure of ether, it may be considered that nine-tenths of the previously dissolved ether and its alkaloid will be recovered in the ministrible solvent. On the other hand the ethereal layer is not wholly free from water which may be expected to take up certain substances not soluble in aphydrous ether, but mactically such traces of impurity are removed on agitating the ether with a limited quantity of water. Similar considerations of solubility apply to treatments with chloroform. but with considerably less force owing to its slight solubility in water and vice-versa, and in the case of petroleum-ether and henzene they have no practical hear-

In making a proximate analysis by means of immasoble solvents, much of the success in practice depends on the care and skill with which the manipulation is conducted. The most convenient apparatus for effecting the treatment consiste of a pear-shaped (fig. 1) or cylindrical (fig. 2) glass esparator, furnished with a tap below and a stopper at the top. The tube below the tap should be ground obliquely so as to prevent loss of hund by immerfect delivery. Supposent that it

ing, as these solvents are almost abso-



he desired to effect the separation of a substance from an aqueous inguid by agitation with other, the former is introduced into the separator, of which it should not occupy more than one-third, acid or alkali added as may be desired, and next a volume of other about equal to that of the aqueous liquid. The stopper is then meersted and the whole thoroughly shaken together for a minute or two, and then set asids As a rule, the mentant will readily separate into two well-defined layers, the lowes of which is aquoons, and the upper ethicsel. Sometimes separation into layers does not occur readily, the Injust remaining apparently homogeneons, forming an emission, or assuming a plathonic sonastency. In such cases, separation may sometimes be induced by thoroughly cooling the contents of the separation. In the cess of ether, the separation may usually be offected by adding an additional quantity of either and ne-eighting, or, when the employment of a sufficient excess of ether is inconvenient or impracheable, the addition of a fow drops of actional following the production of the liquid, will almost invariably cause separation to occur promptic.

The tendancy to form an obstinate enulsion is greatest when the aqueous laquid as alkaline, and is offen very troublesome when chloroform, benzene, or petroleum-ether is substituted for ether. In such cases, the employment of a largor quantity of the solvent sometimes causes separation, but, when admissible, a botten plan is the addition of either. This answers very exceessfully for the isolation of stychmine, which is nearly insoluble in unmixed either, but readily soluble in a mixture of equal uncasures of ether and chloroforim. This efforts in the first plant of the p

Separation having taken place, the aqueous layer should be run off by the tap into another separator, where it can again be agatated with either to insure the complete removal of the body to be dissolved therein. The othereal legular menumer in the first separator should be shaken with a fresh quantity of alkalisated or acadulated water, which is then tapped off as before, and the remaining traces removed by testing the either with a little pure water. This having in turn been run off to the last drop, the ethereal solution can next be removed by the trap, but a preferable plan is to pour it off from the month of the separator, taking care to avoid the draming of any drops of equeous liquid from the sales of the glass.

When anylic alcohol, benzone, or petroleum ether is amployed, the manupulation is the same as that pute desembel, but when chloroform is used, or a mixture containing a considerable proportion of that solvent, the aqueous luqud forms the upper statum, and the chloroformic solution can at once be removed by the tap.

When the volume of fluid treated with the immissible solvent

is very small, the syninge pipette shown in fig 3 may be conveniently substituted for a tapped separator. It is readily constructed by drawing out a test-tube, so as to form a narrow prolongation, the orifice of which should be turned up so as

not to disturb the liquid m which it is immersed. A narrow test-tube fashioned into a handle at the upper part serves as a piston, a short length of india-rubber tubing uniting it to the outer tube, while allowing of easy movement both in a vertical and a horizontal direction.

Another convenient form of separation, devised by W Chattaway, is shown in fig 4 It is practically a small wesh bottle fitting, which is adjusted to the tube or cylinder containing the layers of hund it is desired to separate. It is so arranged that the exittible (B) can be adjusted in height by siding it through

the lower layer.



the india-rubber collar C, so as to bring the turned-up end just above the junction of the two layers. On their blowing through the side-tube (A), the upper stratum is forced up the inner tube, and son be removed, almost to the last drop, without disturbing

The following table shows the behaviour of various classes of organic substances when shaken in acadulated or sikalised solution with immuscible solvents, such as ether, obloroform, amylic alcohol, bensene, and petroleum ether. It must not be supposed, however, that the immuscible solvents can be employed indifferently, but are unaffected by others owing to their limited solvents, but are unaffected by others owing to their limited solubility therein. This is especially the case with the alkaloads and glucosides, and hence the table must merely be regarded as showing their general tendency, then special behaviour with the different solvents being deferred for fuller description lates on.

Table showing the behaviour of Organic Substances with Immiscible Solvents

On agitating the sub-dance with water, actinized with sulpisure cold, and a suitable solvont immiscible therewith (said as ether, etherofers, amytic alcohol, boursus, or petroloum ethers, the following distribution will occur —

hydresurbens, ork, colouring matters, do, which may be agitaling the ligh	LAYER will contain various acids, resins, phenole, glucosides, in there separated in the sealer gan in with sealer gan in the will be	contain carbohydra and ands, organic	agunous Ligum with tes, soluble alkalouds bases, moterds, the her separated by add- langular le sol-
18. THE INSTITUTE OF THE SECOND PROPERTY OF T	18 yell Actually Agency Agency Agency Education and Agency	Is on Dateschae LATER-19 LATER	be ento Accacamo Agracomo Accacamo Agracomo Accacamo Agracomo Agra

The foregoing table meety achiths the general behaviour of the alticulus and other plant-principles on agricing their solutions with manuschle solvents. G Dingendorf, however, his elaborated the following scheme for systematic treatment by manuschle solvents. The statements are made on his subherity, and in some cases (rg, the alleged removal of enchonine from and solutions by chloroform) are of questionable assuracy.

Extraction of Alkaloids and other Principles from Aqueous Liquids by Immiscible Solvents (after Diagondorff)

• The liquid is varied to about 40° or 50° C , rendered distinctly and with distinguishines good, and agitated with an equal reference about the segment of petrolean edders' which is then separated.	e, and separated.	parated	est dissolved chloroform, separated, mnonga, agriated and separated	Aquidus Liquid is agricated with bonzens, and separated	Agreed's Leguin is agreeted with chloroform, and separated.	Acreore Ligure is egitated with a signification of and a separated.	A	
nlphuric acid, and trated.	istre of warm tenzen	h chloroform, and se	etroleum ether to extunded at a	Liquid is agreated w	. Адекова Ілерп	CHLOROPORM ex tracts from the gramomacod	Baretanna Calandan bass Octabadan bass Ontonne Calandan bass Ontonne Calandan Caland	
r 50° C, rendered distinctly saul with dilute gulphs messure of petroloum ether, which is then separated	i Agornous Ligorio is agretated at 40° or 60° O with an equal measure of warm hericans, and separated.	gritated at 40° or 60° C with an equal met		agitated once with po	agituted once with p ether added, then a		honeses sour- from - Zaves- Atropme.	Aydodopermue - Bronne - Bronne - Bronne - Bronne - Brothe - Broth
endered distinctly f petroleum ether,					gumentada gumentad solution —	Anime and methylconne	Pictotine Pagawanna Pintane Passe of cayar emp. purpure, with purpure, we will be purpure, which we will be purpure, which we will be purpure, we will be purpure, which will be purpure, which we will be purpure, which will be	
out 40° or 50° C,					A d	pdang	Canchonne (very doubiful)	Gerssospermue Narceire Papaverne Quebrachne
d is warmed to abo			trom soe genue taked bund Buses-	Berberme (traces)	Colument Paperne (resid- ual portion).	Yon - Baste Prin- cyles— Carvophyllm Cascarilin.	Objects Objects Objects Delphin Delphin Delphin Electrin	
· The hqui		the andulated	Raves-	Non-Basse Prin	Completely) Cops.cum (pun-	Hop Bitter (m completely). Salroyde acid.		

It is evident that the method of Dragendoiff, set forth in the foregoing table, effects some very important differentiations, but for many purposes the process may be simplified with advantage Thus, for instance, if the directions of Diagendorff be adhered to, the aqueous solution will be treated at least seven times (and possibly twice as many) with immiscible solvents before the extraction of morphine with amylic alcohol is attempted. As morphine is not wholly insoluble in the solvents previously used, small quantities, such as are generally met with in toxicological inquiries, are liable to escape detection. Again, agitation of the acidulated liquid with petroleum-ether removes but few active principles, though it is often useful for purifying the liquid from colouring matters, traces of resins, and fatty acids precipitated on acidulating, &c The subsequent treatments with benzene may often be omitted, as the bodies thereby extracted are also dissolved by chloroform They consist of glucosides and other neutral and feebly acid principles, with a few alkaloids of feeble basic character The treatment with petroleum-ether in aumoniacal solution is chiefly of service for the isolation of the volatile bases (conine, micotine, spartcine, &c), and in their absence may often be advantageously omitted. In fact, the three extractions in ammoniacal solution by petioleum-ether, benzene, and chloroform may in many cases be replaced by treatment with chloroform alone, or a mixture of ether and chloroform, which last menstruum possesses the great advantage of separating readily from alkaline

The alkalouts and other principles having been separated into groups by Diagondorff's method, the various solutions may be carefully evaporated and the residues examined for the substances supposed to be present. The special tests suitable for this purpose are described in the sequel.

A tracting alkalode is that of Clars, who for the estimation of affence in reason and of quiume in back, but in capable of many office uptactions. In assaying tea, the powdered leaves are dipid and extracted with ether, the solvent dissolved off, and the renduc extracted with sulphupe and. The acid liquid is filtered, mixed with excess of ignited magnessia, evaporated of tyrness. all, of and the residue pulverised and extracted with boiling ether. For the extraction of quinner, the powdered back is exhausted by water acadulated with sulphure acid, the solution evaporated with excess of magnesias, and the dry residue exhausted with ether.

A Loesch (Year-Book Pharm, 1880, page 60) treats the crude and concentrated alkaloidal solution obtained by suitable

means with three measures of a cold esturated solution of alum, and then adds a slight access of ammona. The liquid contemins the precupitate is evapoated to dryness at 100% and the powdered readure exhausted with a suitable solvent. On evaponating the solution, the alkalond is obtained in colouriess ash-free crystatis. Loesch quotes the following results, as compared with those yielded by Clause magnesia method, and stration by Mayer's solution (page 140):—

/				
	Percentage of Alkaloid Extracted			
Material Employed,	Claus	Mayer	Lossch	
Onchona bark (yellow), { quinno, time banklate, tim	3 176 0 250 1 186 0 500 0 600 2 976 0 146 0 225 0 197 0 825 0 800 0 896	2 570 0 175 1 005 0 895 0 806 0 800 2 800 0 074 0 100 0 725 0 175 0 220	8 250 0 285 1 225 0 825 0 975 8 075 0 175 0 285 0 225 0 875 0 875 0 875	

J U Lloyd (Pharm. Joss., [8], xxi 1144) recommends for the assay of alkaloidal extracts the addition of ferric chloride solution, and then stifficent solid sodium bacerbonate to convert the whole into a paste. This is treated in a procedum mortar with colloroform, which is poured off and the treatment twose repeated The alkaloid is then shaken out from its chloroform solution with dulties and; the latter laund eight attention there is no solution with continues a solution and the same of the th

If a power of the power of the operation it is liberated from its combination by an alkali Should its presence be thus detected or suspected, it may be conveniently isolated by adding excess of lime or barya, and distilling the liquid. The alkaloid can be fixed in the distillate by adding a eight excess of hydrochlona each, and after concentrating the liquid to a small bulk may be liberated by adding a large excess of caustee alkali, and extracted by aritation with either (compare nees 170)

The alkalonds having been obtained in a state of approximate purity by one of the foregoing methods, they may be further treated according to one of the following principles, which may be applied in many cases at an early stage of the process

- a. Fatty and resnows matters and chlorophyll may be removed by sgitating the acidulated solution of the alkaleid with petioleum spirit or ether (Piperine and some glucosides are also extracted.)
- b Colous mg mattles may be removed by agustating the solution with a small quantity of animal charcoal, but this agont must be used very spannely, or the alkalaid may be wholly removed from solution. The alkalaid thus taken up may be recovered by boiling the chancoal with alcohol The absorption of alkalaids by charcoal has been employed for their removal from beer and similar houlds.
- c Many colouring matters, and tannic and various other organic acids, may be removed by treating the neutral solution with lead acctate, and filtering
- d. From sugars, gums, salts, and act active matter symmoly of the alkalouds can be spanted by adding ammons, and agustaing the solution with chloriform or a mixture of allicoroism and other of nesparating the schloroform from the aqueous hquid, which retains the signs, gum and salts, and agritating it with dilute sulphume ead, the alkaloid passes into the acul hquid, while colouring matters, fate, reams, &c, remain in the chloroform
- e The alkaloid may be precipitated with iodised potassium iodide, Mayer's leagent, autic chloride, or platinic chloride, the precipitate being purified by recrystallisation from water, alcohol, or other suitable solvent, and recovered by appropriate means
- By a judicious application of the above pinnaples it is generally an easy matter to isolate alkindos in a nearly puse continuon, or at any rate in such a state as to allow of the special tests being successfully applied. By B P are no "systematic scheme for the proximate analysis of plants detailed "in "Vol. I page 355 et see will also be of servers in the selektion of alkaloids.
- A good example of the separation of alkalous from woody fibre and tannium matters is funnished by the processes for the assay of einchions barks; the separation of alkalous from reasuous, gunnny, and colouring matters is exemplified in the methods for the assay of opinin, while the isolation of stry elinine in terminal termin
- ¹ Made by boiling bone-charcoal with hydrochloric acid, filtering, and thoroughly washing the insoluble residue of carbon

destroyed if the solutions are evaporated at too high a temperature (compare acomitum), (2) that centam alkalouds are extracted by chievatorm and amplic alkabol even from them sendulated solations, (3) that curruine, evitame, morphine, and solatume are nearly or quite insoluble in other or oblordorm, and hence cannot be extrainly extracted by agistant; then alkaline solutions with oither of these solvents, (4) that, whenever possible, the chemical test for the soluted alkalouds should be supplemented by physiological tests, and (5) that, during the process of putrefaction, certain cardavers alkalouds ("ptemmanes") are hable to be formed which simulate some of the reactions of the vegetable bases, but are distinguishable from them by carried acommandion.

CONSTITUTION AND SYNTHESIS OF ALKA-LOIDS.1

Some of the alkaloads of widely different properties present a cumous analogy in their formulae. This resemblance in empirical composition is merely accidental, as is proved in many cases by the products of decomposition. The following are some of the most striking cases of the kind —

1	Atropine,	C17H23NO2	Colchicine,	$C_{22}H_{25}NO_6$
1	Cocame,	C ₁₇ H ₉₁ NO ₄	Narcotine,	CooHo,NO
9	Morphine,	C17H19NO8	Quinine,	CooHan NoO.
	Piperine,	C17H19NO3	Strychnine,	C21H21N2O2
9	Pseudaconitine	C36H49NO12	Piciaconitine,	CalH45NO11
)	Veratrine,	CarHeeNO11	Cevadine,	C32H49NO9

The foregoing coincidences have little theoretical value, as no real insight into the constitution of the alkaloids can be obtained by a consideration of mere empirical formulæ

Some of the most important advances in the synthetical products of alleaded have been due to a study of the products obtained by hydrogenating pyridine and its allies, assisted by a better recognition of the relationship of these bases to each other, and to beneame and other hydrocarbons. Thus, the bases pyridine, quinoline, and aeridine form a senses related to each other exactly in the same way as the hydrocarbons benzene, naphthalene, and anthracene are related (page 39) The stability

Much of the information contained in the test is derived from a lecture by S. P. Sadtlei (*Pharma Jous.*, [3], xx 544), and from the address of A B Prescott to the Chemical Section of the American Association for the Advancement of Science (*Pharma Jous.*, 131, xviii, 520, 541). and the behaviour towards reagents of the corresponding derivatives of benzese and pyridine are exactly analogous, as also is the behaviour on reduction. Just as from benzene hexahydrobenzene can be obtained, so from pyridine hexahydropyridine may be prepared, but far more readily. Similarly from naphibalene and quincline, testrahydro-additive-products are obtainable, while from anthracene and scridine, respectively, dihydro-anthracene and dihydro-actifier (page 128) have been obtained.

Of these hydro-addition-products, one of the best-studied is hexahydropyridine, C_kH_{kl}N, which is identical with the volatale base paperwise, obtamable from paperms, C_kH_{kl}NO₀, the alkaloid of popper, by distillation with alkah, by the action of reducing agents on pyridine, C_kH_kN, or by heating the hydrochloride of pentamethylene-diamine, C_kH_{kl}O(NH₂)_k (page 106)

Another natural plant-base which has been prepared synthetically, and the nature and derivation of white-fire clearly understood, is comes, C,H₂N, the volatile possonous alkaloid of hemiock comms is the darkn-orbatery variety of a normal-propyl-piperidine, C,H₂N, (E,H₂N). To prepare it synthetically, typing then is first converted into a -rall-yl-piperidine, C,H₂N, (C,H₂), which is then reduced in alcohole solution by means of sodium. In this rescion, the chaef product it the optically mactive a-n ormal-propyl-piperidine, which is separated by crystallisation of the textstee into ordinary consist (direct-comine), and a lawo-rotatory comme which closely resembles the other modification.

The optically unactive conne can also be prepared from c on y-rine, or a-conneal-nopyl-nepralme, by treatment with hydrodic such, or from conhydrine, $\hat{\mathbf{C}}_{\mathbf{H}}\mathbf{H}_{\mathbf{J}}\mathbf{N}\hat{\mathbf{O}}_{\mathbf{J}}$, an $\infty\mathbf{y}$ -derivative occurring with connen in henlock \mathbf{A} W H of \mathbf{m} an \mathbf{n} has described three isomence bases, called α , β , and γ -conscense, having the formula \mathbf{m} - \mathbf{n}

Neceting, $C_{\rm L}H_{\rm L}\lambda_{\rm D}$, use vomities auxinom of tobucco, is mother base related to pyraline, and the synthesis of which has been least partially effected. Thus the two known ippyridyls, $C_{\rm L}H_{\rm L}N_{\rm D}$ are the puru- and sast-modifications on reduction, these yueld the corresponding hexahydro-dipyridyls, $C_{\rm L}H_{\rm L}N_{\rm D}$, which are bases called respectively isonicotine.

¹ These two bases bear the same relation to each other, and to the mactive modification that dextro- and lawe-testanc ands bear to racemic acid. Exactly snalogous ethyl-piperidines have been prepared. and nicotidine, isomeric with nicotine. On the atment with oxidining agents nicotine yields nicotining or β -pyridine-carboxylic acid, $C_kH_kNCOOH_l$ a reaction which shows the close relationship of nicotine to the homologues of pyridine

Atropine, C17HoaNOm is another plant-base of which the relationship to pyridine has been very clearly established. Thus when boiled with alkalies atropine is hydrolysed into tropine, C.H., NO, and tiopic acid, C.H., O., while hyoscine, base isomeric with atropine, is similarly split up into tropic acid and pseudotropine Tropine has the constitution of a hydroxyethyl-methyl-tetrahydropyridine, CaHr(CoHaOH)(CHa)N On boiling with acids it loses the elements of water and is converted into tropidine, a liquid base with a comne-like odour, and has been synthesised by Ladenburg by introducing a methyl and hydroxyethyl atom into tetrahydropyridine Tropic acid and tropine reunite to form atropine when their solutions in dilute hydrochloric acid are mixed and evaporated. By substituting other aromatic acids for tropic acid a great variety of bodies can be obtained, which are generically termed tropeines, and one of which, the mandelic acid derivative or homatropine, has proved physiologically important

The pyridine-carboxylic acids (page 110), and their analogues and derivatives, have shown some unexpected relationships to the natural plant-bases The B-pyridine-carboxylic acid (meeting acid) results from the exidation of nicotine and pilocarpine Cinchomeronicacid (a-pyridine-dicarboxylic acid) is produced by the exidation of emchanine, einchonidine, and quinine. One of the pyridine-tricarboxylic acids 18 produced by the oxidation of the cinchona-bases and papaverine with permanganate, while a second results from the oxidation of berberine by nitric acid. The pyridine-carboxylic acids also furnish additive-products analogous to the betaine of beet-juice, and closely related to the natural alkaloids The synthetically produced betains of nicotinic or B-pyridine-carboxylic acid has been shown to be identical with the alkaloid of Trigonella fanuquacum, while the betaine of cinchmogranic acid is identical with a pophyllie a c 1 d. obtained by the oxidation of cotarnine (see narcotine)

The pyridyl-residue, C_gH_kN , is capable of replacing an atom of hydrogen in the molecule of certain acids, the compounds having the same relation to the salts of pyridine that (a,g,) aniline acetate has to acetainlide. *Polocarpaline*, which occurs with pilo-

¹ The same pyridine-carboxyle acid may be obtained by the action of ammonia on coumalinie acid, produced by the action of sulphuric scid on male acid.

earpine in japorandi leaves, is a β -pyridine α - α -1 in 6 thylanid opropionic acid, having probably the constitution \sim NCCH₃, CCH₂- β CH₂- β LH, NCOOH It has been prepared synthetically, and from it, by the action of methyl odde and caustic clickin, followed by silver permanganato, plocapine their has been obtained, and may be regarded as having the following constitution \sim

The setion of ammonia upon certain acids found in the vegetable hingdom has been found to produce boties related to pyridine Thus co in anic soil, $C_iH_iO_{ii}$ derived from meconic acid, is changed by ammonia into an expisionian ceid, while co in a nic acid, $C_iH_iO_{ii}$ from the same source, yields a divexprecionian ceid, come in a nic acid, $C_iH_iO_{ii}$. While solve problems are described in a constant acid, $C_iH_iO_{ii}$ while the alkalonds obsidednms and senguinaries in Chelifonium majus, yields an expyridine-calboxylic acid on textment with ammonia.

Colchaine is another alkaloid the constitution of which is fairly well known. From its reactions shall the products of its decomposition, its evidently the methyl-ester of a cetyl-trimethyl-colchicine acid, (O.OH₂), Co₁, H₄(MH C₉H₂O) CO.OCH₄.

Expensive has the constitution of a methylistinhylitopyndylhydroxypropione and C,H_MMoN GM(GB) GE, GOOD It results, together with methyl alcohol and benzon seal, from the decomposition of cosonie by alkalise Cosasse may be made synthesically by heating execution with benzon anhydride and methyl iodide, and has the following constitution —

C5H6(CH5)N CH(O C7H5O).CH6 COO(CH6).

A series of analogous artificial alkaloids have been prepared by combining eggonine with other acids bosides benzoic

The constitution of the acousts bases is partially known, for they split under the influence of hydrolyming agents into simpler bases and each of the aromanic sense, accountine, prencountine, and igneountine yielding bensole acid, $C_{\rm cH} = 0.00$ H, while pseudocountine gives vorstric or dimethyl-protocate-ahile acid, $C_{\rm cH} = 0.00$ H, $C_{\rm cH} = 0.00$ H. The pseudocountine $C_{\rm cH} = 0.00$ H, C_{\rm

Verative and is also produced, together with $v = 1 n + G_{\rm gal} H_{\rm gN} O_{\rm g}$ by the saponification of verative, while the accompanying base, exactine, $G_{\rm gal} H_{\rm gN} O_{\rm g}$, is converted on hydrolysis into cevine, $G_{\rm gal} H_{\rm gN} O_{\rm g}$, is converted on hydrolysis into cevine, $G_{\rm gal} H_{\rm gN} O_{\rm g}$, and methyl-crotonic acid, $G_{\rm cl} H_{\rm cl} H_{\rm gN} O_{\rm cl}$.

Snapping, C_kH_mNO₂ an alkalond the thocyanate of whuch cause in white mustand seed, is spht by boiling with lenytawater into alinque acid, C_kH_mO₂, and choises, C_kH_mNO₂, a base which is contained in bile and other annual produce as well as in hope and certain other plants. Choine has their been platened synthetically, and has the constitution of hydroxysthyl-trimethyl-ammonium hydroxide, (C_mH,OH)(C/H,N) NOIL.

Theobronuse, $C_1H_2(CH_2)_N V_0$, the alkaled of cocca, and cogions, $C_2H(CH_2)_N V_0$, the alkaled of tea and coffee are respectively the dimethyl and trimethyl derivatives of x anthine, $C_2H_1N_0$, a body occasionally occurring in urmary calculu and produced by the action of intens and on guanine, $C_2H_1N_2O$ (contained in guano), or by treating uric soid,

C5H4N4O3, with sodium amalgam.

Lupning, an alkaloid found in seeds of Lupnius luteus, has the formula $C_{21}H_{28}N_{2}(OH)_{2}$. Argining, $C_{6}H_{14}N_{4}O_{2}$, from the same source (page 178) yields ure a when boiled with baryta water.

The manner in which the oxygen of the natural alkaloude exists is in most cases but little understood. Morphine, $C_{\rm J}H_{\rm L}NO_{\rm S}$ appears, however, to have a phenoho character, and contains two hydroxyl atoms is replaceably by acetyl or beancyl. Codens is a substituted morphine in which one of the hydroxyl atoms is explaced by m eth α xy), $O(B_{\rm L})$ and has been obtained synthetically by heating morphine with methyl iodide. By similar means the second hydroxyl atom can be replaced by methoxyl with formation of methocodesine Theoleum differs from methocodesine by two atoms of hydroxen, thus, —

Morphine, .		C17H17NO OH
Codeine, .	,	CHHINO OCH
Methocodeine,		$C_{17}H_{17}NO \left\{ egin{array}{l} O(CH_8) \\ O(CH_8) \end{array} \right.$
Thebame,		$C_{17}H_{18}NO \begin{cases} O(CH_2) \\ O(CH_3) \end{cases}$

When distilled with zinc-dust, morphine yields phenanthrene $C_{1a}H_{1n}$, and pyridine, C_cH_cN

Narvotne, C. H., NO, is seponfied under estein conditions with formation of meconin, C., H., O., and cotarnine, C., H., O., and cotarnine, C., H., O., and it is latter body when treated with bromine yields dibrom pyridine, C., H., Dr., N. Cotarnine probably contains its oxygen in the forms of OO.OCH., and OCH.

tion it yields apophyllic acid, a body which, when heated under pressure with hydrochloric acid, behaves like the methylester of cinchomeronic acid .--

$$C_6H_3N \left\{ \begin{matrix} \text{CO OH} \\ \text{CO O(CH}_3 \end{matrix} \right\} + \text{HCl} = \text{CH}_3\text{Cl} + C_6H_5N \left\{ \begin{matrix} \text{CO OH} \\ \text{CO OH} \end{matrix} \right. \\ \text{Apophyllic solid} \right. \\ \text{Mothyl chloride} \\ \text{Cmchon-eronic solid} \right.$$

Papaverine, ConHorNO , 1e another opum base, the constitution

of which is probably .—(OCH₂)₂C₂H₄N CH₂C₂H₆(OCH₂)₂.

Brucine, C₂₃H₂₀N₂O₄, when fused with caustic potash yields homologues of pyridine, while strychnine, Con Hoo No Oo, yields

aninoline and indole. C.H.N

Queners, CoaHasNoO, when fused with caustic potash yields methoxy-quinoline, CoHo(O.CHa)N. When subjected to careful oxidation with permanganate or weak chromic acid mixture it at first yields formic acid, CH.O., and a weak base called quitenine, C10H22N2O4. Further oxidation produces three bases, to which Skraup attributes the formulæ C18H18NO0 CoH, NO. and CoH, NO. The first of these has been little studied, the second has been named cinchole up on e. C.H., NO. and the third appears to be identical with kynurine, a base obtained by heating kynurenic soid, a constituent of dog's urine. Besides these bases there are produced cancholeuponic acid, C.H.,NO,;1 quininic acid, C.H.(O.CH.)N COOH; then a pyridine-tricarboxylic acid, C.H.N(CO.OH). and finally the pyridine-dicarboxylic acid known as cinchomeronic acid, C.H.N(COOH), Quintdine and quinicine yield the same products as quinine Cinchonine, C. H. N.O. when similarly subjected to limited oxidation, yields formic acid and cinchotenine, C, H, NoOe, as first products; the latter by further treatment yields ancholeupone, CoH, NO, and this oxidises to oinchoninic acid, CaH, NO, and cincholeuponic acid, CaHisNO, the final products being cinchonic acid, C.H.N COOH (which is a pyridine-carboxylic acid), and cinchomeronic acid (see above) Ourchoudine and emchonicine appear to yield the same products.

The conclusion derivable from the recearches on the constitution of the cinchona bases is that both dumine and cinchonine are derivatives of a hydro-diquinchne, of which probably only one side is hydrogenated. The same unreduced quinoline-residue is common to both alkaloids, with the difference that, while in cinchonine

¹ Cincholeuponic acid probably has the constitution of a methyl-piperidine-dicarboxylic seid -

the readue is quincline itself, in quinine it is a methoxyquinoline. The following formulæ illustrate these deductions.

Quinoline, C_0H_1N $Hydroxyquinoline, C_0H_0(OH)N$ Tetrahydroquinoline, C_0H_1N H $Diquinoline, C_0H_1N$ C_0H_2N C_0H_2N

Cinchonine, . C. H., N C. H., (OH) N CH., Quinne, . C. H. (O CH.) N C. H., (OH) N CH.

Other of the cunchons bases which are known to contein hydraxyl groups are quasammine, C_mH_mN_mQ(OH), and currents C_mH_mN_mQ(OH), The latter sikelend, which is found in Course or Remijia bark, has recently been converted into quanne by heating it to 100°, under pressure, with metalle sodium and a solution of methyl chloride in methyl slochol (Grimaux and Arnold. Comp. Rend. exil 1714).

Although the knowledge of the constitution of the auchons bases is not yet sufficiently perfect to allow of their formation from pyridine or quinoline, it is interesting to note that two distinct basis substances isomeric with quinne have been prepared synthetically, One of these, discovered by C. A. Kohn (Jour Soc Chem Ind, viii. 959), has the constitution of an a-1'-hydroxy-hydroethylene-quinoline,

CoH10(OH)N.CoH4N CoH10(OH).

It was obtained by the action of one molecule of ethylene dibromide on two molecules of α -1-hydroxyhydroquinoline, obtained by reducing hydroxy-quinoline by tin and hydrochlorio soid. It is a weak base, forming small glittering prasms which niet at 283°, and are readily soluble in chloroform and beasses, with difficulty in hot slochol, and insoluble in water. It has weak antipyretic characters

The other synthetical isomer of quinine has been prepared by Wallach and Otto (Annalen, cchii 251) by the reaction of β-nauhthylamine on pinol nitrosochloride.—

C., H., NH., +C., H., O.NOCl = HCl + C., H., N.O.

The product is a basic crystalline substance, melting at 194°-195°, insoluble in water, slightly soluble in hot alcohol, and readily soluble in ether. The solutions, both of the base and its salts, are strongly fluorescent.

Besides the natural plant-bases, a number of bases have been synthetically prepared which have every claim of analogy and character to be ranked with the alkaloids. As instances of these may be mentioned antipyrine, C₁₁H₁₂N₂O (page 32), thalline, C₁₂H₁₂N₁O (page 120), and furture, C₁₂H₁₃N₁O₂.

Y. Meyer has suggested that the formation of the bases and other nitrogenised constituents of plants may be due in some cases to the action of hydroxylamine on aldehydic bodies

It is a curious fact that while the plant-bases and other natural products not unfrequently contain one or more methyl-groups, the ethyl-radical is not met with.

VOLATILE BASES OF VEGETABLE ORIGIN.

Cortain plants contain bases which differ from the ordinary vegetable alkalords, in being volatile, liquid at ordinary or only slightly russed temperatures, and in containing no oxygen. While resembling each other in the above respects, the volatile bases unseen thirth trituther resemblance.

The volatile alkaloids are not numerous, being impated to the following bodies, and a few others which have been but imperfectly investigated.

- Methylamine and Trimethylamine, already described (pages 9, 12).
- b Conine and the associated alkaloids of he in lock.
- c. Lupinine and certain other alkaloids of lupines.
- Nicotine, the volatile alkaloid of tobacco.
- a Psturme, the volatile alkaloid of pituri.
- f. Lobeline, the alkaloid of lobelia.
- g. Spartesne, the alkaloid contained in broom.
- h Spigelme, an alkaloid in Spigelia Marylandica.

Pipersdame, a volatile alkaloid said to exist naturally in pepper as a decomposition-product of piperine, has already been described (page 106)

For the settine-ton at unlead-a-th-1-2 conine in hemlock, and in coti;

nor Amer Ohem Soo, be recommends that a weighted quantity of the substance should be boiled in water accidiated with hydrochloric acid, the residue pressed and washed with water. The solution and washings are evapoisted to one-fourth, and then distilled with slaked line (using a good condenser) When the liquid pussing over is no longer alkaline to litmus, the distillate is exactly neutralised with collected with restricted of the produced condenser and the produced condenser and the second condenser and the condense of the conde

sulphate undasolved, while the sulphate of couns (and other calkainde) pass into the solution. The filtered liquid is evaporated to dryness and the residue skaken three times with caustic potash solution and other, the othered liquid separated and slaken with a known volume of standard sulphure each, the other distilled off or esparated, and the excess of sulphuric and determined by intration. By this process, Loesch found 5 25 per cent of nicotine in tobacco leaves, and 0 06 per cent of comme in the common hearlock plant.

This base has the constitution of an a-normal-propylpipelidine (see page 164)

Comme is the characteristic poisonous alkaloid of hemlock, Consum maculatum. It occurs in all parts of the plant, in combination with organic acids, and in association with the following allied bases —

Base	Formula.	Metting. Pourt, * C	Boling, 'C	Specific Gravity
Ethyl piperidine,	CyHish, or CsHg(CsHs)NH		142-145	{° =0 5874
Comus (Normal-) propyl-piperi- dine).	$C_0 H_{27} N$, or $C_0 H_0 (C_0 H_7) N H$	-25	167-170	{
Methyl-comine,	$C_0H_{20}N$, or $C_0H_0(C_0H_1)N(CH_0)$			{12.5 ± ⇒0 846
Conhydrine,	C ₈ H ₁₇ NO , or C ₅ H ₆ /CHCH CH ₂ CH ₂)NH	120 6	240 (225 at 720	
Pseudo-cophydrine	CaHano, or CaHa(CHa CHaOH CH)NH	100-102	10m) 220-231	

Conine is an only liquid, having a peculiar repulsive odour,

A Confine has been prepared synthetically by the reducing action of soluting another solution of all yl-pyridin, C. High Clip N, stand obtained from e-paroline and paralledgiva. The artificial base thus prepared is sintended in all its properties with the natural elizables, describe that it is optically macrive. But natural alkahori mot on introducing a crystal of the bitarrate of the unstant alkahori mot on very concentrated solution of the bitarrate of the unstant bases, a gradual separation of the beharitate of active commo court, the free base from which exhibit the same optical activity as intuinal comme. The mother-loquid contains a lawy rotatory isomerce have (Laden burg, Ber, 113, 12672).

eth y lalkin, e. C.H. (CHOIL CH. CHI, NML. It presents a descressiblence to trop pine, C.H., NO, both in composition and chemical behaviour, a fact which suggested to A. W. He fin ann the probability that it was the product of the hybridyess of a base aliest to stropine. From the siladime highd left after the distillation of couns and combigation, Ho of man no bisance, by acculation and extraction with ather, caffer a cafe, C.H. (D., a body having the constitution of a dhiphydrey, -in name is a cid.

Conhydrme may be separated from commercial contan, in which it is not unfrequently present, by cooling the hound down to 5° C, filtering through glass wool, and washing the separated crystals of conhydrine with petroleum ether, in which it is but spannight soluble P ac nd oc on hy drin as is a base someone with conhydrine, but probably containing hydroxy-sopropy? (La de n b urg. Ber, xxxi 1871) Conhydrine forms colorites glittering crystals, moderately soluble in water, but very soluble in alcohol and ether the does not react with introus send, has an alkaline reaction, and is a feeble narrotic poison. According to We rith e.in, henlock contains only 6 to 6 parts of conhydring for every 100 of comme

CONICRINES, C. H. N These bases were obtained by A W. Hofmann by the action of oxidising agents on conine, or of dehydrating agents on conhydrine When molecular proportions of conine hydrobromide and bromine are mixed, the bromo-derivatave, C.H., N.HBr Br., is obtained. By the regulated action of caustic soda this yields CaHisNBr, which by treatment with sulphune and is decomposed into hydrobromic and a-consceine. which is a colourless liquid of '893' specific gravity at 15°, boiling at 158°, and alightly soluble in water. In odour it closely resembles comme, but is said to be five or six times as poisonous! It is a tertiary base of strong alkaline reaction, and forms ervstalisable salts. The piciate forms vellow needles melting at 225°. nearly insoluble in cold water, and very slightly soluble in alcohol a-conferme is partially reduced to conine by heating under pressure with fuming hydriodic acid and phosphorus v-coniceine is obtained by decomposing the biomo-derivative CoHigNBr by an alkalı. It is a colourless liquid lighter than water, boiling at 173°, distilling with steam, and said to be twelve times as poisonous as comme! It is only slightly soluble in water, but the solution is strongly alkaline y-conscerne is a secondary base (pages 1.7) yielding crystalline, volatile salts with acids, and a characteristic double salt with stannic chloride, B.H.SnClo which forms large B-conscerne is obtained together with a-conscerne by crystals the action of phosphone anhydride or fuming hydrochloric acid on conhydrine -CsH12NO = CsH15N+H.O. It forms very volstile, colourless needles, melts at 41° and boils at 168°. It is a secondary base of comne-like odour, and is a less active poison than the α -modification

POISONING BY CONINE AND HEMLOCK

Conne is an extremely powerful panalytic poison, which acts on the motor nerves, one drop is a distinctly poisonous dose, while ten drops may be fixtal

The symptoms produced by hemlock and comme are not uniform, and cases of protoning are not numerous Stuppe, come, and slight convulsions have been noticed, while in other cases the chief effect has been passlysse of the muscular system, especially of the logs. The pupils are somewhat dished. After death the lungs are found filled with fitted blood and of a dark colour, and the stomach and intestines somewhat congested. The post-morten unwearances are not characteristic

In toxicological inquiries the viscora and contents of the storacehabould be tracted as desembed under strychnine, the purified extinct being agitated with socla and other instead of ammonia and chlorform. From the ether, the alkaloid may be recovered by allowing the solvent to ovaporate spontaneously in a cool place, or extracted as a sult by agitating the either with dulute hydrochloric and. From the purified salt of comine thus obtained, the free base may be again liberated by adding sook, and recognised by the monty odom of hemlock developed immediately or on warming the inound.

Comme may also be isolated from the viscers by the method used for the assay of hemboch. Otto me one case met with a volatile ptomaine, which was very poisonous, but differed from comine in its reaction with platinic chloride. The seeds of Lepinus Intestes (page 177) contain alkaloids somewhat issembling connic, but which do not yield the characteristic crystalline hybrochloride Chen of the unbediffer ab scalace contain are possessed of poisonous properties, but it does not appear that comine has been proved to be the sactive principle.

¹ Cimanthe a coate, or homlook water-drop wort, as described by A S. Tay lor as one of the most valent of Engish vegetable possons. The leading symptoms produced are raped messashitty, bloated and lavid commensace, convolutive movements, stortows breathing, dilated pupils, and bloody form about the most and nostrules.

Occuta wrosa, water-hom lock or cowbane, produces symptoms similar to the above, including the feaming at the mouth. It is said to contain cioutine.

Sium latifolium and S angustifolium have been mistaken for water cross, with fatal results

Alliusa Cynapum, the lesser hemlock or fool's parsley, appears

Assay of Hemlook and its Preparations.

Comme exists in all parts of the common or spotted h is m lock, Consum maculatum (French, la Cigue, German, der Schiehne) It appears to be most abundant in the fruit, the proportion increasing with the maturity of the seeds. In hemlock leaves, E. K or des found 0.24, and in the fysik 0.49 per cent of alkalod

For the extraction of conne from hemlock, J Schorm (Ber, xiv. 1765) recommends that the fruit should first be swelled by hot water, and then moistened with a strong solution of sodium carbonate. The product is treated with steam, under a pressure of three atmospheree, as long as the distillate has an alkaline reaction, when it is neutralised with hydrochloric acid and evaporated to a weak syrup, which is shaken with twice its measure of strong alcohol and filtered from the precipitated ammonium chloride The filtrate is distilled at 100°, a calculated amount of caustic soda lev added, and the mixture agitated with ether (The residual aqueous liquid developes trimethylamine on prolonged etanding, especially in summer) The ethereal solution deposits large cryetals of conhydrine when strongly cooled. This base is somewhat eparingly soluble in ether, and on distilling the colution passes over with the ether. The conine remaining in the retort is dehydrated with potassium carbonate, and purified by fractional distillation The first 10 per cent, boils between 110° and 168° C., and is very impure. The next 60 per cent, boiling between 168° and 169°, is pure conine, while the next 20 per cent, boiling between 169° and 180°, is impure. The thick dark liquid left in retort contains conhydrine.

A pure product, but somewhat lower yield, so ead to be obtained by exhausting the hemlock fruit with acesta acid, and evaporating the solution to a cyrup in a vacuum. Magnesia is then added, and the mixture agnitated with ether, which extracts the alkaloid.

Many speamens of consum leaves and seed are almost mert from the loss of their volatile active constituent, and hence a method of assay is of considerable importance, and ought to have a place in the Pharmaconoua.

For the determination of the comme and associated alkaloids in bemlock, R. A. Cripps (Pharm. Jour, [3], xvm. 13, 511) recommends the following process—A weight of 5 grammes of the finely-powdered fruit is mixed with an equal weight of sand, and extracted with a mixture of 26 c. of nearly absolute alcohol.

to contain an energotic poison, though this has been disputed by Harley (St. Thomass Hasputal Reports, new series, iv. 63; x 267), and also by Tarret, who believes the erroneous statements respecting it to have arisen from a confusion of the plant with Commun macadatum, which it closely resembles.

15 cc of chloroform, and 10 cc of a saturated solution of dry hydrochloric acid gas in chloroform. The houid is separated from the marc1 and agritated with two separate quantities of 25 cc. of distilled water. The aqueous liquid now contains the conine as hydrochloride. It is shaken once with chloroform, then rendered alkaline with caustic soda, and extracted three times by agitation with chloroform The chloroform is washed by acitation with alkaline water, and is then run into a solution of hydrochloric acid gas in other. This is evaporated in a current of air, and the residue dried at a temperature not exceeding 90° C. The conine hydrochloude obtained should be crystalline, and almost perfectly white From its weight the proportion of conine can be calculated, 1635 of the hydrochloride representing 1270 of the base. If. after weighing the residue, the hydrochloric and be determined by titration with silver nitrate, using potassium chromate as an indicator, the difference will be the weight of alkaloid, and the result should closely correspond with that previously calculated

The foregoing process may be shortened by agitating the weshed chloroforms solution of the conine as hierated by causite soda with water, and gradually adding deemormal hydrochloric scid until a slight and reaction to methyl-orange is developed, which does not disappear on again shaking. Each c. of deemormal add used represents 0 0127 gramme of alkaloid, in terms of conine Petroleum south may be substituted for the chlorofted for the chorofted.

For the estimation of the akkalonds in Tincture of Consum, R art and W right (Pharm. Jour. [8], xxx 857) exports 80 G o. o. of the preparation to a low bulk at 100° C, with 1 c. of normal sulphure acid. The rendual liquid is dubted somewhat at twose shaken with chloroform. It is then rendered alkaline with ammonis, and the heested alkalonds shaken out with chloroform. The chloroform low consumers of ammonis by agitation with water, separated and run into a solution of dry hydrochlore and gas in chloroform, taking one that the orifice of the separator dips below the surface of the said chloroform, which is then evaporated, and the residue dred at 90° and waghted, as recommended by Cripps. The proportion of total alkaloid contained in the intorture of come, as assayed by this process, is from 0.07 to 0.10 per cent. The proportion in the extract ranges from \$\frac{1}{2}\$ on the residue of the control of the con

² The exhaustion should be proved to be complete, by treating the mare with water, and testing the solution with iodine and with Mayer's solution, neither of which should produce more than the faintest turbuity; and the drad mare should give a barely perceptible odour of conine when warmed with causine soda.

Lupine Alkaloids.

From the different species of lupine several alkaloids have been isolated, some of which, at any rate, belong to the class of volatile alkaloids, and in their odour and other characters appear to be related to comme.

LUPENINE, CLITLAND, or CLITLAND, As usolated by G. Braunest from the scale of Lepunes Intern, lupmus no a readily crystalbashle base, incling at 67°5–68° 5, and boiling with some decomposition at 256°–261°. In a stream of hydrogen the distille unchanged at 255°–257°, and is also relatible with steam. Lupmino has a placesant applie-like oldure and an extremely lutter testes, the latter character, extending to its salts. It has paralysing effect on the nevre-entries Lupmine is leave-outstory, easily solidlo in cold water and alcohol, but less solidlo in warm water. From its squorous solition it is separated by excess of austes alkali Lupnine dissolves readily in other, chlorofour, and benness Carrior and the control of the control of

Lupmino is highly eaustie, and is a strong bise, liberting ammonan form its salts and funning with hyllochloris sed is $R(EU)_b$, forms large rhombic crystalls $BH_aP(U)_b$ is crystalline and soluble in water. The amendment $g(R(L))_b$ is $R(R(L))_b$ soluble in water, but resultly in alcohol. The satisfact $R(R(R))_b$ forms rhombic crystals, very soluble in water and alcohol

Metallie sodum dissolves in metad lummie with evolution of hydrogen, forming a sodium derivative, decomposed by water into lumine and sodium hydroxide. When heated with sectic anhydride, lumine yields $C_{\rm B}H_{\rm B}N_{\rm S}(C_{\rm b}H_{\rm b}O)_{\rm R}$ as an oil, misoluble in water and very easily associative.

When lupmine is heated to 150°-180° for ten or breder hours with finning hydrochloire acid, or the hydrochloride to 175° with phosphore anhydride, it yields anhydrotupmine, C_HE_NO, or a lightly oxideable fluid base, smelling like comme. BH₁PtO₃ forms red quadratic tables, easily soluble in water and chits clashol. Denhydrotupmine, C_HE_NN, or satisfable with finning hydrochloire and to 200° G. It is a highly civilable oil, boding at 220°, and forming a so hlor or plast in at in, crystalbang in dark red needles Dephymine, C_HE_NN,O₂ is formed, togethou with anhydrolupmine, by the acknowledge of phosphoric anhydride on lupmine bydrochloride. It is a yellowish, disagreeable smelling oil, boding with some decomnostion at 215°.

Argining, $C_0H_{14}N_4\tilde{O}_{2\nu}$ is contained in the seeds of *L. luteus* which have germinated in the dark. It forms crystalline salts, evolves nitrogen with introus seed, and yields ures when boiled with baryta-water.

LUPINIDINE, $C_2 \coprod_{10} N$, is a base found by Baumers in tha y balue or time that y balue and proper in the property of the surface of the surface and feebly possonous, producing symptoms like those of course Lupinidus forms a crystalline hydrate, B.H.O. very modulule in water. The saiks are crystalline to a control of the surface of

LUPANINE, C15Ho, NoO, 18 an alkaloid obtained by M Hagen (Liebig's Annalen, coxxx 367, Jour Ohem Soc, 1 163) from the seeds of the blue lupine, Lupinus angustifolius, which are stated not to contain lupinius or lupinidine. It is described as a pale yellow, honey-like syrup, with green fluorescence, intensely bitter taste, and an unpleasant odour like that of hemlock Lupanine does not boil at 290°, even under the reduced pressure of 130 mm It has a strong alkaline reaction, attacks the skin, expels ammonia from its salts, and forms with hydrochloric acid white fumes of the hydrochloride With excess of cold water, lupanine forms a turbid solution, from which the base is almost entirely separated on heating. It dissolves with difficulty in cold alcohol, but readily in ether, chloroform, and petroleum spirit Lupanine hydrochloride, BHCl+2aq, forms hygroscopic, quadratic crystals, melting at 127°, and soluble in alcohol but not in ether. BHoPtCla is not distinctly crystalline BHAuCl, forms golden needles, insoluble in water, alcohol, or ether From solutions of its salts, lupanine is precipitated by caustic potash and soda, but not by ammonia.

From the seeds of Lapraus albus, Campan usolated a poisonous liquid alkaloid, boiling at 210°-218° From the same source Betelli obtained a crystallisable base

According to O Ke'lln ex (Biad Cents, x, 97) lupuse seeds can be deprived of the whole of their bitter constituents, and rendered much more palitable and wholesome, by soaking them in water for twenty-four hours, steaming them for one hour, and then weaking them for two days Kuhn has shown that the substances which cause lupuse schoses are destroyed by steaming

Nicotine. Nicotin $C_{10}H_1 M_2$, or $C_8H_1 N$, $C_8H_2 N$. Nicotine has the constitution of a he x h hydro-dipyridyl (see page 164). It is the poisonous base principle of tobacco, in which it exists combined with malie and citric acids (compare page 184), in proportions varying within very wide limits.

First modifies a colourless, only fluid of 1011 specific gravity at 15° C. On prolonged exposure to air it becomes yellow, and eventually resmoid. It has a sharp caustic taste, is intensely poisonous, and has a strong and unpleasant odour, recalling that of tobacco. Nicotine bolis at about 250° C, with partial decompositions of the process of the pr

position, but it distals readily with the vapour of water or alcohol, and volatilises to a notable extent at the ordinary temperature. Nincotine absorbs moisture from the air, and evolves heat when mixed with water, diminution in volume simultaneously occurring.

Skalweit (Ber, xiv 1809) has given the following figures showing the specific gravity of mixtures of micotine and water. His results point to the existence of a hydrate of micotine.

Freotone	Water	Specific Gravity at 15° C
100 100 100 100 100 100 100 100 100	0 6 10 20 80 40 50 00	1 011 1 017 1 024 1 030 1 034 1 057 1 040 1 038 1 088

Nucotine has a powerful levo-rolatory action on polarised light, the value of S, m. 20 per sent aqueous solution being, according to Pribram, $-161^{\circ}55$ The rotation diminishes rapidly but irregularly by further dultuon. Thus for a 4 per cent solution the value S, in $-77^{\circ}03$, while below this strength an increase is observed, S, being $-79^{\circ}03$, while below this strength an increase is observed, S, being $-79^{\circ}03$ for a solution of 0.8326 specific gravity. The rotation is affected by time, not reaching its maximum for 48 hours (Eq. , xx 1340)

The aqueous solution of nicotine is powerfully alkaline in reaction. The motions is partially separated by addition of excess of causic potash or sold (compare pyridine). Nicotine in aqueous solution, and in the absence of other free base, can be determined by titration with standard lead and methyl-ownnee.

Nuosane forms two classes of salts. The monanci salts are stable and neutral to thums and methyl-rousage, but the ducated salts have an acid reaction. Most of the salts of months cyrathline with difficulty. The acid Land Lat, $\Omega_{\rm ph} H_{\rm s} N_{\rm c} (L_{\rm H} Q_0)_{\rm ph} + 2\alpha_{\rm f}$, as an exception, and forms handsome thrife when other is added to its achieble is altical to its

DETECTION AND DETERMINATION OF NICOTINE

Alcohol dissolves meeting in all proportions, and on evaporating

When water us added to solution of mostone contaming less than 20 per cust of base, the mixtus becomes tribid and clears only on long standing. On instanç to 40° the liquid clears rapidly, but becomes again turbid when cooled or further heated to 50°. Heterosof 0° and 0° the turbidity unmanate to multimess, which disappears when the highest is cooled below 50°. At 70° the nontine separates in part as an oily layer.

or distilling the solution the alkaloid is found chiefly in the first fractions. It is extincted from its aqueous alkaline solutions by agitation with ethics, chihorform, bennene, anylice alcohol, or peticoleum apints, and may be received from the solvent by separating and agitating with dilute sends. If oxalic and be employed, the resultant solution may be evaporated to dryness and treated with alcohol, which dissolves the meetine exalits while leaving any ammonium existing undessolved. After again removing the alcohol by evaporation, the montime may be theretaed from the warm luquid by adding crosses of caustice solds, when the characteristic tobacco-like small of motume will be observed, and the although can be obtained pure by distilling the highed with water, or egistating it with either and allowing the separated solvent to synaptica socializations.

Treated with nitric acid, meetine yields a thick reddish liquid. Sulphuric acid produces no change in the cold, but a brown colour

is developed on heatang.

On dissolving mostuse in dulus hydrochlore soid, and adding platine chloride, sicotize chicopations, O.g., N.M., P.-Club, is thrown down as a spanngly soluble, yellowish, crystaline common The prequisate is soluble in hot wasts, especially in presence of free hydrochlore soid Addition of alcohol increases the delacacy of the test, and the formation of the prospitate is much facilitated by stirring the liquid. A m m on n gives a smill seathon, but on n is given a smill seathon, but on n is relief and the common of the prospitate of the control of the common of the prospitate is much facilitated by stirring the liquid. A m m on n gives a smill seathon, but on n in a yield m or necessitate with alsatine chloride.

Prorie acid, if added in excess to solution of meetine, throws down cotine picrate as an amorphous yellow precipitate, which rapi changes to a mass of crystalline tufts, even in presence of

for organic matter.

thme is precipitated by Mayer's reagent (page 138) from v. lute solutions; and, by operating in strongly and liquids, 27. It's by obtained very good quantitative results. The formula of the precipitate is Challagh and I c. of the reagent represents 00020 gramme of motions.

On adding mercurne chloride to a solution of modune a white regretalizes promputes is produced, soluble in dulte hydrochloric or aceins and This is the most characteristic reaction of modules as summar proputates, nearly modulies in acein acid. This is the most characteristic reaction of module in acein acid. Many other alkaloids are precupitated, but the compounds are almost invastably amorphous. This is the case with the precupitate produced by counce, which is almost the only alkaloid which will distill over with notions on boiling the solution with a slight excess of caustic soda. Am mon in a, however, behaves like motions, and must, if necessary, be separated before applying the

test Ammonia is sharply distinguished from mootine, conine, and lobeline by adding a solution of notine in nodicie of potessium to the slightly accelerated solution of the base. Ammonia produces no change, but with either of the vegetable alkaloids a brown or brownish and piccipitate will result. Todine solution will detect 1 of mootine in 250,000, and is the most delenter regards thowy for the alkaloid.

Solutions of montme are not preeputated by chromates, ferrocyanides, ferroyandes or throcyanites, nor by gallic and With gallotamno and an aqueous solution of motome yields a white, amorphous preeputate, which readily dissolves on cautious addition of hydrochloric acid, but is again preeputated by further addition of acid, and is then usedolble even in a large excess. Tannate of motome is readily soluble also in acetic and nitro acids, but is not repreeputated on adding an excess

A variety of processes have been devised for the determination of mootine in tobacco and its preparations. The problem is complicated by the presence of ammonium salts, by the difficulty of completely extracting micotine from aqueous liquids by agitation with immissible solvents, and by the tendency to form an emulsion when these are used, owing to the presence of pectinous matter, The methods proposed have been reviewed by J. Biel (Pharm Zest Russ, xxvii 3, Analyst, xin. 97), who iscommends the following process, which is a modification of that proposed by Kissling .- 100 grammes of powdered tobacco-leaves, or 10 to 20 grammes of extract of tobacco, are mixed with slaked lime and distilled in a current of steam until the condensed steam is no longer alkaline. The distillate, which will measure about 1 litre, is rendered faintly and with dilute sulphuic acid, evaporated to 50 c c., made alkaline with caustic sods, and acutated six times with other, using 20 cc each time. Biel then distils off the greater part of ether slowly, adds excess of decomormal sulphuric acid, and titrates back with decinormal soda, using resolic acid as an indicator. The object in distilling off the ether is to get rid of any traces of ammonia which may be present, but it is difficult to do thus without risking the volatilisation of some of the nicotine It is preferable to titrate the unconcentrated ethereal solution by gradually adding decinormal sulphuric acid, using methyl-orange as an indicator, and agritating between each addition. Each cc of deconormal acid neutralised represents 0 0162 gramme of micotino The results will be high if ammonia be present, and in such case the neutralised aqueous liquid should be separated from the ether, and evaporated to dryness at 100°. The residue is weighed and treated with absolute alcohol, which will dissolve the sulphate of meetine, while any ammonium sulphate will be left mealuble, and its weight oan to deducted from the weight of the nursed sulphates previously found, the difference being the sulphate of nucebane. The result may be confirmed by adding placed, phthalient to the alcoholic solution of nucebane and its titrating with decunormal alkali, which will sear just as if the sulphure and were meanulant.

From conine, meetine is distinguished by its odour, by being heavier instead of lighter than water, and by the reactions with hydrochloric acid gas, mercuric chloride, argentic intrate, platinic chloride, and pience acid (see above, and page 181).

Poisoning by Nicotine and Tobacco

Nicotine is one of the most violent poisons known. Only a few instances are on record of poisoning of the human subject by the pute alkinoid, but the effects of tohacoo, which owes its poisonous properties entirely to mootine, are well known. I impute solutions of necotine and infusions of tohacoo are employed as insecticacles

"The usual effects of a possonous does of tobacco, when taken into the stousach, are confusion in the head, paleness of the countenance, vertigo, nauses, severe retelling and vonuting, heat in the stomach, great anxiety, a sense of surking at the pit of the stomach with extreme prestration, trendling of the limbs, and sometimes vrolent purging. The pulse is small, feeble, and almost improceptible, the respiration difficult, and the skim cold and clammy, the pupils are generally dilated, but sometimes contracted, and the vision is usually more or less impaired. Death is often preceded by convisions and paralysis" (T. G. Wormley, Murcochemutro of Poisson).

In toxicological investigations, incotine may be isolated from the vacers in the same namers as connic (toges 170, 176). An alternative method is to digest the suspected matters with water acadulated with acosts send, and treat the filtered liquid with accessed flead scatals. The liquid is again filtered, the lead removed from the filtrate by passing sulphurested hydrogen, and the clear solution treated with caustic soda, separated from any precipitate, and distilled, when a fluid having the odour and exhibiting the reactions of nuctine will be obtained. Any supposed mootine which may be isolated should be tested by placing it on the tonges of a young rability or small bird, when themose, paralysis, and

¹ When tobacco as an &k ol, the grater part of the mecture a convention into pradits and other progenous compounds, and the earbr decomposition of the mecture is sometimes assetted, but Meisens appears to have fully proved the presence of unchanged monotone to tobacco smokes ma proportion equal to about one-seventh of that present in the original tobacco (compass equal to about one-seventh of that present in the original tobacco (compass equal to about one-seventh of that present in the original tobacco (compass equal to about one-seventh or that present in the original tobacco (compass equal to about one-seventh or the present in the original tobacco (compass equal tobacco).

convulsions will rapidly ensue Nicotine appears to be unchanged by putrefaction, and hence may be detected in the tissues long after death

. Товасоо (French, le Tabac, German, der Tabah).

Tobacco is the dried leaf of Nicotianum Tabacum and allied

According to S W. Johnson, a good crop of tobacco, yielding 1260 lbs. of dry leaf and 1110 lbs of dry stalk, removes from the soil the following constituents in lbs. per acro.—

Constituents.	onts. Leaves		Total	
SO _b , FyO _s , CaO, MgO, KgO, NgO, NgO, Ngo, Ngo, Nim of Ash Constituents, Nikrogen,	74 78 78 17 71 5 206	\$ 15 16 2 47 10 95i . 88	17 221 88 19 118 15 3011 82	

As the stalks are returned to the land, tobacco is not a very exhausting crop, but requires abundant manuring, since the period of growth does not exceed three months. Hence, tyo may be advantageously sown as soon as the tobacco is off, and ploughed in as a green oney when utilityation for tobacco commences.

Besides cellulose, albumunoid compounds, pectic acid, guarconstants a peculiar volatid, crystalline principle called nicotinini or tobacco-campho, to which the formula CAHANAO, has been attributed Tobacco also contains the volatile alkaloid nicotine, which is apparently peculiar to the genus The base exists in combination with mails catif, but the presence of citrates, acctates, and oxalates has also been established. Fresh tobacco-leaves (sepecully the midrie) contain a notable proportion of nitrates, but these saits are easiled to disappear during the process of fermentation to which manufactured tobacco is subjected. This fermentation has for

¹ The genus Nicoteanz contains more than 70 species N. Tabaccum yields the tobacco of Havana, Cubs. France, Holland, Belgium, &c. N. rusticzz furmshee East Indian tobacco, and the kinds known as Letakia and Tulkish tobacco. Tum be k to r Persan tobacco is the product of N. Persan.

² From 100 grammee of dried tobacco leaves, Goupel obtained from 3 to 4 grammes of acid malate of ammonum, J. Takayams (Chem. News. 1.

constituents, and the formation of "ferment oils," which probably contribute to the aroma, especially when saccharme matter,

800) obtained the following percentage results by the analysis of Japanese tabanco --

1	Nagato.	Shtmozuki	Settzu.	Osuzni
Water, Ash, Nicotine, Acade acid Oxalic acid, Malio acid, Citric acid, Pectac acid,	8 41	10 01	7 68	13 18
	15 78	8 45	20 71	9 30
	2 45	8 02	8 92	1 89
	0 05	0 04	0 01	0 98
	trace	0 27	0 25	trace
	0 79	1 02	1 83	2 08
	0 62	0 69	0 92	0 89
	1 24	5 84	7 62	2 35

In the above analyses, the nectine was extracted by ammoniscal ether, the solvent distilled off, and the mootine in the residue determined by titration For the acetic acid, the powdered tobacco was moistened with water and tartane acid, and distilled in a current of steam, the acetic acid hang determined in the distillate. For the fixed organic scids, 10 grammes of the sample was moretened with sulphune acid in the quantity requierte to combine with the bases (as indicated by the carbonates in the ash), and exhausted with ether From the ethereal solution the acids were extracted by a small quantity of water, the separated aqueous liquid rendered alkaline with eramonic acidulated with acetic and, and the oxalic acid precimitated by adding esloum acetate. To the filtrate, a driute solution of lead acetate was gradually added, until a test quantity of 1 c.o of the supernatant liquid gave, on further addition of lead acetate, a precapitate which was completely soluble in a few drops of acetic acid. The liquid was then filtered, and the precupitate of lead citrate washed with water containing a little lead acetate and acetic acid, and then with alcohol, the washings being kept separate The citric acid was deduced from the weight of lead oxido left on igniting the precipitate. From the filtrate, the make acid was prompitated by excess of lead acetate colution, and its amount deduced from the weight of lead oxide left on ignition. The washings from the precipitate of lead citrate were boiled to expel alcohol and treated with excess of lead acetate, the precipitate being regarded as a mixture of lead citrate and malate in equal proportions (compare Vol I. page 484)

The yelf-t end was determined by schaming 10 grammes of bolecon with controlled spint continuing one-fourth of its volume of concentrated hydrochlore soil. The roades was washed with spirit till the hydrochlore soil was whelly removed, and then treated with a solution of a known weight of ammonium oxisite, by which the pecte soil was dissolved. After digestrate of two hours at ST, the liquid was filtered, the results wested, and the filtrate shifted to I little. As aliquot part of the was preceptated by colours and the state of the

liquorice or alcohol is added during the maceration to which the tobacco is subjected 1

As sold by the farmers, the tobacco-leaves contain about 30 per cent of wator. When the fresh leaf is simply dried, the product is yellow, the brown colour of commercial tobacco being due to the regulated fermentation already alluded to. The unmanufactured tobacco imported into England is converted into ioll or spun tobacco, cut tobacco, and cigars, the refuse being used for making snuff. In the manufacture of roll-tobacco, the leaves are moistened with water, spun into various sizes of twist, made up into iolls, and piessed. The liquid or juice which exudes is used as a sheep-dip. Cut tobacco is made by moistening the leaves, cutting them to the required size, and drying on plates, or it may be made into cakes first, and afterwards cut The Excise regulations prohibit the use of any foreign matter in manufacturing tobacco, besides water and a little oil Hence, except in the proportion of water, which is not allowed to exceed 35 per cent (as estimated by drying at 100° C), there is no tangible difference between manufactured tobacco and the dried leaves imported. The proportion of micotine in tobacco does not appear to be an index of the quality

J. Clark (Jour Soc Chem. Ind., in 554) has published the percentages of ash yielded by the ignition of twenty-one authentic

			In 100 Parts of the Dry Substance.			
			Total Ash	Soluble Ash, "Alkaline Salts"	Sand.	
Whole Leaf — Highest, Lowest, Average, Limins — Highest, Lowest, Average, Middle —	:	:	39 80 * 18 79 30 52 81 67 * 12 47 19 21	11 87 2 40† 6 47 8 99 1 95† 4 98	12 82 * 0 18 2 68 14 41 + 0 09 2 86	
Highest, Lowest, Average,			30 87 * 15 14 21 92	90 (II 4 03 11 41	4-91 * 0 12 1 15	

^{*} Paraguay Tobacco

samples of representative to bacco-leaves. The table is an abstract of his figures, which in all cases refer to the leaf dried at 100° C

[†] Ohinese Tobacco

¹ Schizomycetes occur in fermented tobacco in large numbers, but the number of species is very limited. Thial experiments by E. Suchsland, with foreign ferments on German tobacco-leaves, yielded a tobacco not recognisable as of German organ.

As the composition of the lamins and of the stem or midrib of the leaf differ materially, these were carefully separated before analysis

E Quajat (Their Centr, 1880, p. 345) found the ash of fourteen samples of dry tobacco (neulump both superur and common knuls) to ange from \$1.03 per cent in a Bassano sample to \$171 in Virgamm and 1678 per cent in Turksh. He considers that the quality of tobacco varies inversely with the ash, but Nessler recognises no relation between the two

I r by and Ca be 11 (Chem Nees, xxx 117) have published the figures obtained by the analysus of ax typical samples of Vaginan tobacco All were in the leaf state, free from stalk, but staming the multib No 1 was light yellow tobacco, "coal-caused varappers" for eggers, No. 2, light yellow, "fine smoking" tobacco, No. 3 was medium brown colour, "sweet fillen" for eggers, No. 4 was dath; "Austaina and Italian egger wampers," No. 5, dark "Engiha shipping," and No. 6, dark, "exported to France for south". These samples when an-druck yelloble for samples when an-druck yelloble for the first property of the sample when an-druck yelloble for the first property of the first property

	No 1	No 2.	No 8.	No 4	No 8	No 0
MOISTURE, per cent , ASH, total, per cent on tobacco, ,, Soluble in HCl, per 100 of ash, ,, Sandand charcoal, ,, ,, ,, Carbon diovide, ,, ,,	7 91 11 80 70 71 6 30 23 99	1 00 15 39 69 17 14 09 28 14	11 07 18 52 60 93 16 98 22 09	9 93 16 31 84 40 7 92 7 08	13 74 18 18 64 83 8 82 20 65	9 71 16 00 66 66 8 97 24 37

Deducting the sand, carbon, and carbon dioxide, as also the small proportions of alumna and feare oxide found in the portion of the sah soluble in said, the "pure sah" of the tobacco was calculated The total introgen was determined by the absolute method of D um as, and the mootine by Ma ye'r's solution, with the following results, expressed on 100 parts of tobacco dried at 100° C :—

	No 1	No 2	No 8	No 4	No 5	No 8	wge.
Prue sell, Total nitrogen, Nicotine, Nitrogen in forms other than nicotine, Percentage of tutal nitro gen present as nicotine,	8 p4 3 18 3 32 2 61 18 2	9 29 2 63 3 59 2 01 23 6	12 84 3 72 5 28 2 81 24 5	14 84 5 76 7 09 4 54 21 3	13 89 5 33 6 20 4 20 20 1	11 05 5 25 8 85 8 73 98 9	11 64 4 89 5 79 8 83 22 8

The following table shows the average proportions of nitrogen

and ash, and the composition of the latter in tobacco from various sources .--

	Virginia.	Krnydory	NEW ENGLAND	Runophan (including Turkish)
Observer, Number of spessmens con tributing to average, Maindeen, per cent, "Purs ash," per cent,	Irby & Cabell 6 4 82 11 64	Peter 80 12.83	S W Johnson 12 4 24 16 56	E Wolff 18
Percentage composition of ash- SiOp Cl, SO _h , PyO _b , RyO _c Na _y O _c Na _y O _c MgO _c	1 72 2 81 5 19 3 30 35 58 278 37 60 19 72	278 874 421 499 87 57 2 10 35 31 9 35	0 84 9 86 6 58 8 56 34 96 1 99 84 48 8 21	10 28 4 92 4 30 8 21 18 01 4 20 48 51 11 48

Will and Fresenus (Ann Chem. Phanm., 1877) have recorded the issults of them analyses of the sah of a number of samples of Hunganan tobacco, and Schlossin, kxxx 148) the proportions of potash, lime, magness, sulbates, and chlords in the sah of tobacco grown on different soils. The proportion and composition of the sah of English tobacco has been investigated by A Wingham (Joseph Seo Chem Ind., v. 78, 400), of Indian and Burmess tobaccos by R R om an is (Chem News, 1812, 248), and of varous kinds of tobacco grown in Japan by J Takayama (Chem News, 1 301), and Fesca and Ima (Jour. Seo Chem Ind., vin. 759)

The combustibility of tobacco is profoundly affected by the proportion and nature of the universal constituents, especially the calcium and potassium, and the forms of combination in which these metals occur. The ash of the more combustible tobaccos is comparatively rich in potassium carbonate, showing the presence of a large proportion of organic salts of potassium in the original tobacco, while the ash of tobacco of inferior burning quality contams a larger proportion of sulphates or chlorides, and hence proportionately less alkaline carbonates According to Schloesing and Nessler tobacco burns best when it contains a considerable proportion of potassium malate, which is a natural constituent of the leaf, but the effect may be imitated, and a slow burning tobacco improved, by the addition of potassium acetate or other organic salt of potassium, while the combustibility may be diminished by addition of sulphate of ------According to E R Durrwell the white tobacco 18 due to the presence of a large proportion of alkaline salts, which swall up as the tobacco burns, and tear the fibes, thereby indees, complete combuston. Stiphates rather favour proper combuston, while interies are psyndicial. Chlordes are regarded by most observers as objectionable, and hence should be absent from fertities interied for empirication to tobacco-cross.

A Mayer (Land Yersuche-Stat, xxxvii 127, Jour Chem. Soc, Jivii. 148b) has investigated the influence of various unstances employed in 0.5 per cent, solution on the combustibility of ordinary filter-paper. Organic substances of the most different kinds were found favourable to combustion with fiame and to diminish the power of glowing, while inorganic substances usually had the opposite effect ²

From his experiments with filter-paper, Mayer concludes that the more ash tobacco yields, and especially the more potential carbonate (representing organic aslts of potessum in the tobacco), the better the obacco will burn, while much calcum phosphate, sulphate, or chloride is held to be prejudical. The alkalimity of the ash is a better measure of combustibility than the proposition of chlorine. Mayer gives the following figures obtained by the partial analysis of tobacco of different unfilties from Sumatra

Товдосо	CHLORINE	Total POTASH	ALKALINITY 85 K ₂ CO ₃	ASH	NITROGEN
Good, Sufficiently good (light) ash), Sufficiently good, Sufficiently good (grey) ash), Bad,	07	50 68 66 79 46	49 68 65 41 05	20 5 20 8 22 5 17 7 18 5	27 32 20 33 26

- 1.0. Cantoni (Bud Gente, 1879, p. 813) found that mixtues of the initial initial had most effects, seguridar typou of growth of the tokenoc when it is a small control in the state of the initial initial had been chierale and Fireway and the same that the print in weight being scatally ingles who as manner was practically such as amonous sulphate or actions chierale was added. The had was almost totally uncombratible whom the plant had been manners with gream, but this produced by meaning the state of the same than the statement that othercale are objectionable in tokeno manners, and states that their use increases the proportion of chlorine in the leaves from 0.21 to 0.50 per cent.
- ² The salts found most favourable for glowing wee the alkalms attacks, sulphates, and carbonatos, alkalms organic salts, and potassium chloride. Sodium salts had less effect than potassium salts, and calcium and magnesium salts much less still. Paper treated with potassium salts, magnesium sulphate, or sodium carbonate pare a white salt. Glorieds were found wither to favour glowing.

According to J. M. van Bemmelin (Lossel Vesscha-Schot, xxxvu 409, Junu 3Ass), Junu 3Ass), to be see which burns budly other contains an excess of chlome and sulphure acid over the potash, or eise the anomat of potassum, compared with that of chlorme and sulphure acid, as low, owing to the potash barng partially replaced by soid. Lawse of the best quality contain hild or no soda, not much chlorme or sulphure said, but a large proportion of organic saids of potassum, acidium, and magnesium. Too much fat or albimin in the tolacco neutralises the good effect of organic saids of potassum, and it is important the total culturing the casing of the tobacco in the sulficentity decomposed during the casing of the tobacco in the said the ratio of Co₂ CH+SO₂ is not best than 2 1.

According to M Φ or, tobacco which burns hadly can be made to burn well by sleeping it for twenty-four house in a 0.5 per cent. of potassum acotate on intents. In this way soluble organic matter and alkalime chlorides are extracted, while the selfs is two-unable to glowing are taken up. By steeping in a 0.5 per cent solution of calculum acetate, the more immonities this choice, which can not otherwise only be used for small, can be made to burn well, and yield a nerfectly white sah

The mode of existence of the nitrogen in tobacco has been investigated by Fesca and Imai (Jow Soc Chem. Ind., vii 759), who have published the following among other interesting

analytical data .1___

	Highest	Lowest	Average
	Percentage	Per centage	of 8 Samples
In structured bodesers standing and structured by the structure of the st	1 91 13 21 14 64 19 4 73 14 44 16 50 1 69 0 67 3 61 4 69 4 09 41 3 409 48 6	1 02 8 29 10 88 0 84 5 06 5 14 10 94 11 17 1 29 0 82 0 89 8 63	1 48 10 46 12 52 0 44 3 54 8 97 12 12 14 10 0 48 2 58 3 16 88 7 29 2 88 1

¹ Fesca and Imai deduce the following conclusions from their researches — The quantity of meetine may be considered as bearing the same relation to tobacco as the percentage of alcohol does to spirituous liquors; but as yet a

high percentage of meeting has not been shown to be an indication of the good quality of tobacco. Nitire and should not be found in well-fermented tobacces Ammonia determinations are frequently too high, as they include some auncle-nitrogen. 0-1 per cent or so of aunmonia does not seem to lower the quality of the tobacco. The albuminoids in a tobacco afford no indication of quality unless the proportion of amides is simultaneously coundered. The amide-nitrogen represents for the most part harmless, or, perhaps, even beneficial, nitrogenous compounds. It is possible that a further study of these bedies and their decompositions will reveal the presence of bodies evencising a direct influence on the quality of tobacce. Anyway, the conversion of albuminoids into amides is one of the most important results of the fsimentation Oldmany fat determinations, or rather extracts, are of no 1150 in tobacco analysis. Carbohydrates should not be present in well-formented. tobacco, but a study of the changes they undergo would doubtless be of great valus in connection with tobacco Only considerable differences in the amount of the various constituents of tobacco can give any conclusive indication of the quality of a tobacco. Very had tobaccos always contain much albuminoid matter sulphune and, chlorine, and large quantities of mineral acids, with small proportions of ample nitrogen, potash, &c By the present methods of analysis it is easier to recognise a bad tobacco than one of good quality Bases, particularly potash and lime, in medium quantity, are favourable to the good quality, and especially the combustibility, of tobacco. An excess of enthst of these bases over a liberal mean percentage is neither a sign of good quality not combustibility, and only an exceptionally low percentage of orthor of them can be regarded with containty as a bad sign. Very high magnetia is projudicial to the combustibility Mineral ands in large quantities indicate both bad combustibility and quality, but only a very high proportion of au individual acid can be safely considered a decidedly had indication. The combustibility is influenced to the greatest extent by the quantity of sulphune acid prescut, and in a diminishing degree by the percentage of chlorine, phosphoric acid, and silica in the tobacco. The percentage of soluble curbonates appears to have no unportant influence on the quality and combustibility of tobacco, the influence of the total quantity of embenates in the ash is much greater, but even in this there is a maximum beyond which the percentage of carbonic anhydride in the ash cannot be regarded as indicating increase of combinstability The relation of carbonates to the mineral acids is a much more important factor, a large preponderance of the former being a favourable sign High basicity of ash is an excellent indication of good combustibility, especially when not due either entirely, or to a great extent, to magnesia or iron.

lead acetato still contains a sugar-like body, which he did not attempt to isolate, and which had little or no optical activity, but which yielded alcohol on fermentation with yeast, in amount corresponding to an average of 7 per cent of sugar Eastes and Ince (Phasm Jour, [3], xvi 682) found a small percentage (2 5 to 5 3) of a fermentable saccharoid matter, not removable by lead acetate, in the extract of tumbeki or Persian tobacco (Nucotrana Persica) The nucotine in this product ranges from 2 to nearly 6 per cent., and the ash from 22 to 28 per cent

H. Muller (Bied Centr, 1886, p 409, Jour. Chem Soc., 1 904) states that fermented tobacco contains, as a rule, little or no starch, and no sugar The whole of the starch commonly disappears during the first few days of the drying. The sugar thus formed is often converted into water and carbon dioxide, and this change seems to be complete in leaves quickly dried. The last trace of sugar disappears when fermentation sets in, while any residual staich does not appear to be altered.

From the analyses already quoted, it is evident that the proportion of micotine in tobacco varies considerably 1 According to Schloesing (Chem. Gazette, v. 43) dried French tobacco contams from 5 to 8 per cent, of the alkaloid, Virginia and Kantucky, 6 to 7 per cent., while Maryland and Havana tobaccos contain only about 2 per cent, and oldmary snuff about the same proportion L Ricciardi (Ber. xi. 1385) to some extent confirms those results, for he found the micotine in twenty specimens of tobacco, grown in Italy under various conditions, to range from 5 99 per cent in a Virginian variety to 1 62 in Havana tobacco.

Tobacco Smoke varies in character according to the proportion of air admitted during combustion, oxidation being necessarily more perfect in the case of a cigar than when the tobacco is smoked in a pipe In the latter case, a portion of the condensible products is deposited in the liquid state. Tobacco-smoke consists in part of ... permanent gases, the proportions of carbon dioxide and carbon monoxide in which have been determined by G Krause Vohl found sulphuretted hydrogen and hydrocyanic acid, and from 0 7 to 2.8 grammes of ammonia for 100 of tobacco smoked Vohl and Eulenberg (Arch Pharm, [2], cxlvs. 130) experimented on the smoke of strong tobacco, burnt both in pipes and in the form of cigars. The smoke was first aspirated through a solution of .

? According to Ad. Mayer a liberal amount of heat and light, together with sufficient moisture in a rich soil, will not only cause a lexurious development of tobacco plants, but give a large increase in the percentage of mootine. while the other organic constituents of the plant are not much affected by climatic conditions

caustic potash, and then through dilute sulphure acid. The alkala absorbed action dixxide, sulphuretted hydrogen, hydrograms, formic, acetic, propionic, butyric and valence acids, placed and ecococts, the presence of captoc, caprilles, and succinic acide could not be accustaged conclusively. The acid absorbed ammonis, pyridnic, $G_{\rm LH}[H_N]$, and all the homologues of the sense to viridina, $G_{\rm LH}[H_N]$, michiave, in addition to the above, earbon monoxide, methane, and several hydrocarbons of the acciylence series were detected. Pyridine visual series in the smoke from pupes, while collidate was the prominent base in cigar—smoke.

Vol.1 and Kulenberg conclude that the motions of tobacco is completely decomposed dump the process of smoking, and that the intense action of tobacco-smoldo on the nervous system is due to the presence of bases of the pyridine series. These is no violattial some observers have metaken these bases for micotine, but that some observers have metaken these bases for micotine, but that some observers have metaken these bases for micotine, but the les on a "experiments (Dunl Polyt Jour. Xiv. 121) appear to be conclusive as to the presence of micotine, which he isolated in a condition fit for analysis and to the amount of about 33 grammes for 4½ kitigammes of tobacco smoked, or about oneseventh of the unmarky nucestally mescale!

Tolocco Emicar varies greatly in strength, and should always be assayed for the proportion of mootime. A good extract is and to contain about 7 per cent of the alkaloid. The following analyses by E G siefler (Jon. Soc Ohem Hid., viii. 426), of tolocco extract of 40° Baumé, impostes a wide difference in its charactes, according as it is prepared from the leaves or midnles of the tolocco.

	Liquid	Mineral Matter,	Containing E ₂ CO ₃	Organio Matter	Containing Niedtme
Extract from leaves,	35 2	15 6	5.0	50 83	81 .
Extract from midribs,	52.8	221	7 73	48 40	1 86

Sough is manufactured from refuse-tobacco, such as siems, tobaccomails, and sweepings. These are mostened with water, subjected to a process of feinmentation during six or eight weeks, then ground, mixed with alkaline salls as pieservatives, and flavoured as desired in the United Kingdom, nothing is allowed to be added to snuff

¹ M clasna* conclusion has been endorsed by R Kussling (Ding Polyt, Jour., ochir 64), who has collected and wraves when the observations of previous investigators. He considers V ohl's conclusion as to the non-custome or honotome in tobacco-smoke to be due to that chemists having overlooked the fact that the silizated as decomposed by waim caustic pokesh, a reaction which, if a fact, has estatishy not met with general recognition.

but the chlorides, sulphates, and carbonates of notassium and sodium, and the carbonate of ammonium, and any smult which contains a greater proportion of these salts than 26 per cent on the dry snuff, including the salts natural to the tobacco, is liable to forfeiture and a penalty of £50 As the proportion of alkaline salts in tobacco-ash varies considerably, it is important that the manufacturer should know the amount present, in order that he may compound a snuff of uniform composition, and not exceed the legal limit. Of the salls allowed to be added to snuff, common salt and the carbonates of notassium and ammonium are those most commonly used. In addition, most snuff contains from 25 to 45 per cent of water, and sometimes a considerable quantity of sand, the proportion, according to J. Clark (Jour Soc Chem Ind., in 554), averaging 5 per cent on the day snuff, but ranging from 0.5 to over 10 per cent, and in one case exceeding 30 per cent A large number of gross and more or less apocryphal adulterants of smuff have been recorded. Among these the sulphides of arsenic. mercury and antimony, chromate of lead, bichromate of potassium, sulphates of copper and iron, alum, lamp-black, avory-black, cream of tartas, sed ochre, buck-dust, and various organic matters find a place As snuff is neither a "drug" nor an article of food, it is not liable to examination under the Adultoration Acts, and the Excise systematically ignore sophistications which do not affect the revenue Hence, authentic information respecting the present adulterations of snuff is very limited.

Piturine. C12H16N2

Pituime, the volatile alkaloid of pituri, was regaided by Petit as identical with micotine, but its distinct individuality has been established by Liveisidge (Pharm Jour, [3], xi 815)

In its chemical characters and physiological effects pitting presents the olosest sessibilities to mostine, but is distinguished from that base by its reaction with P alm's test. When gently warmed with hydrochloric and of 112 specific gravity, in c of 1:no times voicel, and on addition of a little storag untrie and the colour changes to a deep orange. Pittings when thus treated does not change solum at all, but when further heat is applied it turns yellow

Pitume s distinguished from conine by its aquicous solution not becoming turbed on heating, or by the addition of chlorinewater, from aniline it is distinguished by its negative reaction with solution of bleaching powder, and from picoline by because somewhat densor than water. From pyridine, pitumo differs by

¹ Pitun consists of the dried leaves of *Dubousia Hopwoodus*, a shrub growing in Australia. It centains from 1 to 2½ per cent, of the alkaloid

LOBELINE 195

giving a precipitate with cupile sulphate insoluble in excess of the base.

When pittene is treated in othereal solution with iodius (compare spiration) the highid becomes hownish tool and turbid, and after a short time deposits yellowish red needles, leaving a yellow mother-diptor. The crystals melt at about 110° G, and dissolive in adouble with hownish led colour. This solution leaves indistinct needles and only drops on evaporation; it breaten in the cold with caustic soda, an indeform-like odour is evolved, whereas the iodius-compound of incline is said to reproduce uncoins when similarly treated.

Lobeline is the active principle of Lobelia influta, or Indian tobacco, a plant which has received extensive application by unauthorised practitioners, and is also an official drug ¹

LOBELINE exists in lobolia in combination with a vegetable acid In presence of certain other constituents of the plant the alkaloid is extremely unstable, being rapidly decomposed on heating an aqueous, or even an alcoholic, infusion of lobelia. In presence of acetic acid the base is more stable, and was obtained by J W. and C G Lloyd (Pharm Jour, [3], xvii 1038, xviii 135) as a colourless, odourless, amorphous substance, permanent in the air, only slightly soluble in water, but readily soluble in alcohol, other, chloroform, benzene, carbon disulphide, &c In the nure state lobeline is not hygroscopic, and is but slowly changed on exposure to air Lobeline turns red with sulphuric acid, vellow with natric acid, and is precipitated by all the general alkaloidal reagents. The salts, which have not been obtained crystallised, are readily soluble in water, alcohol, and ether. They are described as most violent emetics, a single drop of a tolerably strong solution producing immediate emesis, without disagreeable after-symptoms The dust is as irritating as veratrine to the nose and air-passages.

¹ The canne dired hee's constitutes the official ding, but the dural see of a followin as also hargely used. The rot of to Locken application was employed before \(\tilde{L} \) righted was known to medicane, but the root of the latter species does not appear to have been used. According to \(\tilde{L} \) W, and C \(\tilde{L} \) I o'd, all parts of lobelia contain the alkaloid, which, however, is most readily obtained from the seeds.

The dust of the plant produces a panful sometors when mhaled all parts of the heth and seed produce on actal butung senation on the tongue, and a sharp, tobacco-like impression on the threat and fancer. Lobdin contracts the pupil, and acts as an expectorist in small doses and a memor in slegtred to to 20 gramp. In poisonous quentities it also like mechan, and kills by paralyang the respuration. Several fatal cases of poisoning by lobelia are on record.

INTLATIN was obtained by J. W and C. G. Lloyd in large colourless, doubtes crystals, melling at 225°, insoluble in molecular or glycerin, but soluble in alcohol, ethor, chloroform, benzeue, carbon dissulphie, and the oil of boles, &c. Infiatus is a necessprinciple, and appears to have no therapeutic value. The lobelaus is of Enders is considered by the Lloyds to be a mixture of infiatus, rosm, lobeline, and the fixed out which lobels contains in the uncorstant of about 30 use cent

No liquid volatile alkaloid could be obtained by Messis Lloyd from lobelia, by distilling the herb with water, other with or with out the addition of caustic alkali, and they considered the supposed volatile base to have been probably a mixture of lobeline, inflatin, and volatile of

On the other hand, Paschkis and Smita (Monath, xx. 1811, Jour. Soc. Oken. Ind., ix 761) have obtained a volatile alkadual from Lobela suflata, by extracting the leaves with water activated with aceit acid, rendering the concentrated solution alkalina, and aguisting with ether. On distilling off the solvent the alkadual is obtained as a viscous oil, with an odour stonce resembling that of honey and tobacco. It is purified by solution in didite hydrochloric acid, and re-extracted by alkali and ether! After distilling off the other the base is drade with caustic potach, and distilled in a current of hydrogen. On warming the alkadual co-obtained with a 10 per cent solution of caustic potach, and gradually adding a 4 per cent, aqueous solution of potassium permanganate, be n zor a card is formed, and can be extracted by filtering off the precipitated oxide of manganese, and agitating the acciditated solution with the excellibrated solution or the form.

The sulphate of the above volatile alkaloid, if prepared from lobelia seeds, is obtained in yellow, very hygroscopic granules. When prepared from the leaves, it forms a yellowish white powder, less hygroscopic than the salt from the former source

According to Drese n lobeline is the only medicinally active principle contained in Lobelia uplata. S. Nunez (Brit Maz.) Jour., 1889, 1059) considers it greatly superior to the galenical preparations of lobelia, and recommends it in the treatment of spasmodical asthma and bronchiteal dyspin.

¹ Up to this point the process of Psychias and Smits is substantially the same as that of the Lioyd Bros for the preparation of the non-veitable alkaland of lobels. Sie bort, by the same process, has recently obtained, both from the bets and seeds of lobels, a pale yallow skilmin symp, the crystallised hydrochloride and chloroplatinate of which indicated the formula C₁₈H₂₈NO₂ for the free alkalon.

SPARTEINE 197

Sparteine C₁₅H₂₆N₂

This sikaloid is obtained by H ou dé and Labord e (Pharm. Jons. [3], xv. 543) by exhausting in a deplacement-apparise with proof-spirit the coarsely-powdered leaves and brunches of be to on (Sgatum seeparamy). The product is filtered, dishel under reduced pressure, the resedue dissolved in tartarc each, the luquid filtered to remove a greenish deposit containing chloryland seeparam, Ca_BH₂O₃₀, the filtrate tendered alkalime by potassium carbonales, and agitated several times with either the chieves solution is shaken with tartarc each, and the separated alkalime and extracted with chieve, which on evaporation leaves the alkaloid; the yield being about 0.3 per cent of the blant used.

Spatane is a colourless, oily hquid, bohing at 287° at the ordernary pressure, on at 180° at 20 mm. It has a somewhat pump, pyridue-like odour, a very bitter isste, and on exposure to air times brown and thick. It is soluble in alcohol, either, and chlorothen, but insoluble in petroleum either. It is solution in alcohol (24° per cont) has a senselin totation of —14° for the sorium ray.

Spartame is a well-defined base, uniting with ands to form crystalisable salts, and having the constitution of a tertany diamine. The sulphate forms large, transparent, very soluble rhombehedia, a solution of which gives with caustic alkales and ammonia a white precipitate insoluble in excess. Cadimum nodizing gives a white curify precipitate, and sodium phosphomolybidiate a white precipitate, dissolving on heating the hquid. Platinum chloride yields a yellow precipitate of BHLPCQ+ Eaq, very insoluble in cold water and alcohol, but crystalliang from hydrochloric and in rhombuc prams. Spartame gives no coloration with concentrated mineral ands

When oxidised with potassium permanganate, sparteine yields a ¹ Administered in doses of 0.1 gramme, Sparteine sulphate is stated (G. Seq.

The state of the s

According to De Rymon, spartous causes trame, dilation of the pupils, inco-ordination of movements, and convulsions alternately tonic and clonic Schroff found that a diop of spartone introduced into a labbit's mouth occasioned spasms of the numeles of the spine and limbs and paralysis of the latter, slowing of the respiration and bent, and death in six minutes

The effects of spartone have been compared to those of comme, but they do not explain the value of broom as a director medicine

amall quantity of a volatile (apparently fatty) acid, tegether with a non-volatile pyridine-carbexylic acid, which on distillation with lime yields pyridine Heated in sealed tubes with fluming hydriodic acid, spatione yields mothyl iodide and a base containing C., MLN.

According to Be in he imer, en gradually adding 3 parts of sortine dissolved in ether to an ethereal solution of 1 part of sportens a black preceptate is formed, which, when separated, washed with other, and dissolved in boding alcohol, crystalines on ceoling in beautiful green needles containing Ci₁₁T₂₀N₀I₃I₃. This body is miscibile in cell water or alcohol, but dissolves in either fiquid when heated. It is miscibile in either promiser in the air, and yields free sparteins when heated with caustic alkal (compare "Pitturne," grage 195). Bromme acts strongly on sparteine at the ordinary temperature, even when largely diluted with ether, forming an undefined remnous mass.

According to Grandvaland Valser, when a drop of ammenum sulphydrate is placed on a watch-glass, and a trace of spartenne or one of its salts added to it, a permanent orange-red coloration is immediately produced.

Spigelime is the active principle of Spigelia Morplandica, or "pink" not." as detailed by W. L. Din dle by by discipling the root with mulk of lines it was volatile, gave with todine as howenish-red precupitate, and with Mayora' reagent a with erystalline precipitate seluble in alcohol and ether, and differing from most similar precipitates by being soluble in dishes acid, Spigeline is said by Si a bler to be little, precipitated by tannin, and seluble in water and alcohol, but not in either (i). Pink"-root is effect used as a veinifuge, and possesses possoness properties albed to those of gelsemium, depressing the action of the heart and of respitation, and in large doses causing less of muscular power (Pactitioner, July 1887; Asiaer Chem Jour, 138). It produces strubiamus, dilatation of the pupils, and temporary loss of sight, with some drowness but not narcotsum. A fluid extract of spigals root is official in the US. Pharmacopeas.

ACONITE BASES,1

The different species of Aconstum contain alkaloids of a cleselyallied character, but which differ from each other in their chemical

¹ The subjects of this section are discussed at greater length and in more detail than their intrinsic importance seems to waitant, but it appears desircomposition and physiological activity. The characteristic seomito alkaloids are perhaps the most violent poisons known, but certain spaces of aconite contain simply harmless, butter principles. All parts of the plant contain the poison, but the not is richest unlarked. If any poiston of a poisonous aconite plant be cleaved, it will be found to have a taste which may be at first bitterish sweet, but there a time becomes such and burning, causing a persistent sense of tinging and numbness of the gums and tongue, which effect less for some time and is hughly characteristic.

For medicanal use, the German and United States Pharmacop on the absorbers root of Acountum Mapshine (Wolf's-bane or Monk's-hood). The extract of acounts of the British Pharmacop one as prepared from the fresh leaves and flowering tops of A Napshine ("gatheted when about one-third of the flowers are expanded, from plants cultivated more than the state of the description of which points to an impuse product), the linim out and the tineture are directed to be prepared from the carefully-dried mot of the same plant ("collected in winder or early spring before the leaves have appeared, from plants cultivated in Bittan or unpried in a dried state from Germany "). The French God ex authories the use of the leaf and root of both A. Napshine and A freez, possibly

able to present the chemistry of the acousts bases in a more complete four than has been done since the publication of Aldei Wight's classical researches ending in 1880. The author is indebted to Dr C R Alder Wight for regusal and correction of the article.

³ The root of Magadhu us from 2 to 4 mehrs long, and of an uregister content form I is much sinveiled longituhnality, and is more observed with the sears and bases of broken rootlets. Externally, it is cofference, but the transverse section as whitain, and extubuts a contrai cofficient, a character which distingualises account root from house-admit, which it temotrity resembles, and for which it has been statilly mustaken. The detailed is temotry resembles, and for which it has been statilly mustaken. The detailed Regges (Pharm Jose, 18] vm 912, Okeans and Dragges (Pharm Jose, 18] vm 912, Okeans and Dragges (Pharm Jose), 18] vm 912, okeans and Builting from the Structure of A. heterophyllium and Jaqueses accounts have been described municity by W as over (Charm Jone, 18), x 3011; xx 140)

Noire/instanting the impotance, in the case of such a dug as scottle, of unitering stretch to the directions of the Phart na cop pra is, it is sketed on the high amount-not as most with in common or signarily of German or Japaness origin, the former being gathered indoscrimately from plants which may vary as widely in "properties as A developphilium (non-posinous) and A. ferce (highly posinous), and certainly do vary as much as A. Negellus (intensely posinous) and parentalized (non-posinous).

owing to the widely-spread, but apparently mistaken, impression that the alkaloid known as Morson's acousting is prepared from the latter species (compare foot-note on page 216).

The note of accunic plants are not only the nehest in total alkaloidal contents, but the alkaloida extracted from the root of A Napelius were found by C R Alder Wright to contain a much larger proportion of the crystalline base acon it in a than the alkaloids from the other parts of the plant (stem, leaves, and flowers).

The vanous natural alkalouds of the acountes are, broadly speaking, characterists of particular species of the plant. Thus a con it in me is the poculiar alkaloud of A. Nagellus, pseudacon it in each five half of A. Nagellus, pseudacon it in ed. A. Nagellus, and it is includy include that the traces of pseudaconine found by A.I.d. or Wright in the alkalouds from A. Nagellus, and, conversely, the trace of acontinue discheded in the bases from A. Force, were due to unsuspected admixtures of other species of acontin in the patries of roots which professedly came from one spicess only. Thus, twenty-mine varieties of A. Nagellus have been described, and some of these calculated and differences that only an expert could distinguish thom from nearly allied species. The true A. Nagellus flowers in May, and appears to be peculiar in this respect; it is impossible even for a skilled botanist to distinguish the plant by its leaves alone (E. M. Holl me s. Plant Morr. Jour. (51, xu. 756)*

The roots of at least two species of Japanese accomite come in the United States, viz. Accomiten Feeberic and A senicination. The latter species has been described as poisonous, but, according to V Coble nt. 3, the root, although it contains an although security devoid of the taughing and numbing tasts of A Napellus The physiological expariments of Bartho low on the root of A. Fischer: indicate that this plant increases the number and droot of the cardine pulsations, instead of reidoung the heart's action to M. A Napellus. These and other results show that juncomitine and preparations of the Japanese root should by no mean be substituted for A. Napellus for internal administration (Pharm. Jour., [8], xri 545).

Besides the eminently poisonous alkaloids, acontine, pseudacontine and japaconitine, characteristic respectively of Acontium Napellus, A. ferox, and A Fischeri, other species of aconte

Mandelin, by the examination of A. Napellus alkaloids of various degrees of purity, was not able to detect pseudacontine; and Jurgens also failed.

The root of Imperators Ostruthsum, or mostorwort, has been not with as an adulterant of acounte. It resembles acounte tubers in shape, but has an aromatic odour and pungent taste, and the transverse section exhibits numerous off cells arranged in several circles.

contain alkaloids which appear in some cases to be highly possible outs, and in other cases hursless, bitter tomes. This the biland of Aconiton paracelatam (which was the official acounts of the London and Dublik Planracopporas of 1839) as a mert, bitter principle, not improbably identical with the pic aconities including by T. B. Grove a from a part proposal to be those of A. Nagellus. The root of A. hate ophyllum contains a nort probable that similar bases occur in other species. Lyacondition and myoctomas are physiologically active alkaloids solitated for A lyacotonium. Some species of according appears to the contain an unsolitate base shaving distinct a narrow to provide the narrow of the property of the contain an unsolitate base shaving distinct narrows for present such such as the contain an unsolitate base shaving distinct narrows present see to according papear to contain an unsolitate base shaving distinct narrows prepietates.

The following table shows the chief sources of the aconite alkaloids and their derived bases. The root is the part of the plant referred to in each case.

PLANT	Saponifiable Bases	Basic Products of Suponification	Unsaponuflable Alkaloids
Aconstum Napellus Monk's-hood Wolf's- bans (blue flowers)	Aconitine Amorphous base Picraconitino (ex ceptionally pre sent)	Aconine ? Pieraconine	Amorphous unnamed base
	Pasudaconitine (very amall quantity, if at all)	Proudaconine	
Aconstum Feroz Indian aconste Nopaul aconste Himalayarcot "Bikh" or "Bish"	Pseudaconitine Amorphous base (7). Aconitine (in very small quantity, if at all)	Pecudaconine ? Acomne.	Amorphous unnamed base
Aconitum Anthora (yellow ish or white flowers)	Pseudaconitine (7)	Paendaconine	
Aconstum Fischeri. Japanese aconite	Japaconitine. Amorphous base (f)	Japaconine	Amorphous unnamed
Acontum Unematum.	Bitter inactive alka	1	
Aconstum Pansoulatum	loid Patraconitine (?)	Picraconine (7)	
Asonitum Lycoctonum (yel low flowers)	Lyaconitine	Lyaconine (Lycocko-	
low Howers)	Myoctonine.	Lyaconine (Lycocto-	
Acontton Heterophyllum (blue or dirty yellow flowers, with purple veins) Atis or Atees root.	1	t mme)	Atisine.

Constitution and Characters of the Aconite Bases.

Much of the earlier work on the alkaloids of the acoustes is of little value, owing to the readiness with which the bases undergo decomposition, and the consequent failure of the observers to obtain them in a pure state,

The following table shows the leading properties of the better known of the acouste bases.

Name.	Synonyme and Sources	Formula	Melting- Point,	Appearance and Characters	Physiological Effect
Acoultine,	Napaconitine Crystallised acomtine From A Napellus	C ₅₀ H ₄₅ NO ₁₂	188	Crystallizable both in free state and as saits Alialoid dex- tre retatory Salts lave retatory.	Intensely personeus
Anhydro aconi- tine,	Apoacouitine	C38H43NO11	188	Small coherent crys	As poisonous as aconitina,
Aconino,	Saponification of acoustine	C16H41 NO11	180	Amorphous, forms amorphoussalts Rs ducos Feilling's solu- tion	Bitter; moder ataly poison ous.
Pseudacopitino,	Acraconitine, napelliue, feraconitine. From A	OMPH SHOP	105	Base and saits crystal lise with difficulty Saponifiable	Intensely poisonous.
Pseudaconine,	Saponification of pseudaconi- tino	CHH41NO	100	Amorphous; forms amorphous salts Does not reduce Fahling's solution	Bitter, slightly poisonous,
Japaconitine,	Crystallino alkaloid of Japanese acquite root	C ₆₁ H ₈₈ N ₂ O ₂₁	184-180	Ctystallisable, forms crystallisable salts. Sapoulflable	Very poison ous, closely resembles accritise
Japaconine,	Saponification of inpaconi tino	C28H42NO9		Amorphous, forms amorphous salts Re- duces Fehling's solu- tion.	Closely resembles aconine
Pieraconitine,	Doubtful; per haps the fu active alka- loid of A ponseulatum	Cal Has NO 10	above 100	Bass crystallises with difficulty but salts easily Saponinablo	Bitter, vo poisonous
Piersconine,	Saponification of ploraconi	CHHUNO!		Amorphous	Bitter, no poisonous
Lyaconitino, .	From root of	C27H24N2O8	112-114	Amorphous, deviro rotatory Saponifi able	Poisonous
Myoctonine, .	With lyacom tine, in A	C46H51N2O12	144	Amorphous, dextro rotatosy Saponin able	Bitter, para lytic polson.
Lysconino,	Lycoctomus Saponifica- tion of lya conitine	CzrHayNuO7	46	Crystallisable, dextro	Poisonous
Acolyctine,	With lyaconine			White powder	Paralytto
Attsino,	From root of A hetero phyllum	C46H74N2O4	85	Forms crystalline baloid salts	Posson. Bitto, no possonous.

It is not probable that either the foregoing list or that on last page includes all the distinct alkaloidal principles of the aconites. The so-called "amorphous alkaloids" have been very imperfectly examined, owing to the difficulty of obtaining them in a condition of purity. Of those which have been partially examined, considerable uncentainty exists as to how far they are natural considerable uncentainty exists as to how far they are natural considerable uncentainty exists as to how far they are natural constituents of the original plant, and how far formed by polymenisation or other changes during the process of extraction. T and II. S in it to better the process of extraction. T and II. S in the obtained to not help face, justed of the looks of A. Negellus an altalond which appeared to be narcotine, and which they termed a con-cultine T be occurrence of this beek has the best confirmed, but it is noteworthy that there is a relation to been confirmed, but it is noteworthy that there is a relation in the constitution of narcotine and pseudomentaion, for which the former yields me conin, C_0H_0 , or opianic act, C_0H_0 , C_0 , or appendication, the latter gives of un at hyl-protocatech unce actd. The following formulæ show the constitution of the two instruments.

$$C_{\delta}H_{2} \begin{cases} O \ CH_{3} \\ O \ CH_{3} \\ CO \ OH \\ CO \ H \end{cases} \qquad C_{0}H_{2} \begin{cases} O \ CH_{3} \\ O \ CH_{3} \\ O \ CH_{3} \\ O \ CH_{3} \end{cases}$$

The researches of O R Alder Wright have demonstanted that the crystallashle alkalous of Acoustan Napelius, A perox, and A Fuscher (Japanese acousts) are alkyl asits or esters, either of benzone acid taself or of a derivative of this acid Thus, when heated with alkales or mineral acids, or to some extent when heated with water alone, acid to the orgadizine bases undergoes seponification, with foundation of benzon acid, or a derivative thereof, together with a new amorphous base of far less physiological activity than the crystalline alkaloud from which it is derived.

The following table shows the composition of the natural crystalisable alkaloids of the group, and the products of their signoification. The formule of accounts and account are those attainfued to these besses by Dunstan and Inco (Jour. Chem. Soc., in 271), and show H_s more in the molecule than the formule of Alder Wright for the same alkaloids.

³ The statement made in the text sequires qualification Proteomities a sagenfishle alkaloid, but in not possenses. It forms seathly replicable salts, but the free base has not been obtained caystallised. Atmins, again, as itself amerphous, but forms crystallisable labeled salts, and as not known to be separafiable. Ingenentiae and mycotoniae have not been obtained crystallised, but are sapenifiable and yield crystallisable salts.

	PRODUCTS OF HYDROLYSIS			
CRYSTALLINE BASE	Amorphous Base	Antd		
Aconitine,	Aconiuo,	Benzoic acid,		
Cg3H45NO ₁₃	C ₂₀ H ₄₂ NO ₁₁	C7H ₆ O ₂		
Picrasonitine,	Pioraconine,	Benzoic and,		
C ₃₂ H ₄₅ NO ₂₆	CalHalNOp	C ₇ H ₆ O ₂		
Japaconitine,	Japaconine,	Benzoic sold,		
CasH ₅₈ N ₂ O ₂₁	20 MH 41 NO 10	2CyHgO2		
Pseudacomitas, C ₂₆ H ₄₉ NO ₁₂	Proudeconine, C ₂ H ₄ NO ₉	Versirfo sold (dimethyl protocatechnic soul), C ₀ H ₁₀ O ₄		

Lyacontine, the amorphous alkaloid of A lycoctonum, also yields an acid and one or more bases on sepondication (of which one, lyocotonine, leadily crystallises), but it doubtful if the reaction can be expressed by any simple formula (see page 223). The amorphous alkaloid mycotomie, from the same source, yields bensoic acid on sepondication, together with the crystalline base lycoctonue, and other products

The saponification of the crystalline aconite bases occurs with a near approach to quantitative accuracy, at least so far as the production of the acid products is concerned, the basic product usually undergoing some further change with formation of a resmous substance The reaction is best effected by boiling the alkaloid with alcoholic caustic soda for some time, under a reflux condenser If the product be then acidulated with hydrochloric acid, and agreated with other, the acid products of the saponification are dissolved. On separating the ethereal solution, and shaking it with soda, the benzoic and veratric acids are dissolved, while resmous matter remains in the ether On again acidulating the separated alkaline bound the acids are liberated. and may be dissolved out by agitation with ether After allowing the washed ethereal solution to evaporate spontaneously, and drying the residue over sulphuric acid, the acids may be weighed, or, where only one is present, the amount may be ascertained by tatrating the ethereal solution with standard alkali and phenolphthalem A method adapted for the assay of very small quantities of the acomite bases, and based on this principle, is described on page 234 After weighing, the melting-point of the acid may be ascertained Benzoic acid melts at 121° C., and may be separated from veratric acid (page 218) by prolonged distillation with water, when only the former body passes over. The distillate may be rendered alkaline, concentrated to a small bulk, acidulated, and the beazon and extracted with ether, and recovered by evaporation of the solution. The veratire and may be similarly recovered from the liquid left in the retort

The following table shows the proportions of carbon and hydrogen contained in the crystallisable acounte bases, together with the percentage of gold contained in their aurochlorides (direct at 100°), and the proportion of acid yielded on saponification.—

Alkslord	Pormuls	Carbon	Hydrogen.	Gold in Anto- abloride	And by Saponifi cation	NaHO required for Saponist cation.
Aconitino, Pacudaconitine, Picraconitme, Japaconitmo,	C ₅₅ H ₄₅ NO ₁₂	01 20	6 95	19 98	18 92	6 20
	C ₅₆ H ₄₅ NO ₁₂	02 88	7 18	19 10	26 49	5 82
	C ₅₁ H ₄₅ NO ₁₀	02 95	7 61	21 07	20 60	6 77
	C ₅₅ H ₅₅ N ₂ O ₂₁	63 67	7 07	20 89	19 60	6 48

When the hydrolysas of the natural aconate bases is effected by heating with concentrated mineral excis, or even by water stone under high piessure, the saponification is pieceded or accompanied by the removal of the elements of water and a formation of the so-called "apo-bases," preferably called anhydro-bases. The following table shows the relation of the apo- or anhydro-bases the parent alkaloids, and exhibits the constitutional formulas of the former —

The suhydro-bases are best prepared by beating the pursue alkalouis to 100° for sur to ten hours with a esturated egapous solution of tartaire seed. On rendering the higher distance with sodium bicarbonate, and elaking with other, the anhydro-base is dissolved, and may be obtained in crystals on evaporation of the etherest solution.

In their physiological effects, the anhydro-bases resemble the alkaloids from which they are derived. Thus "apo-acontine," or anhydro-acontino, is extremely poisonous, while anhydro-aconine is nearly inactive.

Japaconitine, the natural alkaloid of Japanese aconite root, undergoes no further change when heated with tartaric acid, for it has the constitution of a sesquianhydro-derivative.—

$$C_{36}H_{36}NO_{7} \begin{cases} 0 & \text{CO.C}_{6}H_{6} \\ 0 & \text{CO.C}_{6}H_{6} \\ 0 \end{cases}$$

The hydrogen of the OH-groups of the anhydro-bases is capable of replacement by organic acid-acides! Thus when pseudocontine is beated to 100° for some hours with a large excess of gland sector acid, it loses the elements of wate, but the sub-you-base formed is then further acted on with formation of a c t y l - an h y d ropseu d a contine, which is a base cyreallaism (like the parent alkabad and its anhylro-base) with 1H₂O, foruing a crystalline mittale and gold sait, and yriding section and dimethyl-protocatechure acids on sepomfestion with alkables. The same product is obtained if acits snipydride be used in place of acets acid, while, if benzies snipydride be substituted the corresponding to a r so y l-d e r r a f i r v is produced. When acountine is hested with bemome anhydride it yelski, in a smillar manner, b e n z o y l-a n h y d r o a c o n i t i ne, a product which is spiparently identical with the robust mend by the action of benzies anhydride committee.

207

Japacontine is converted by benzoic anhydride into a derivative containing four benzoyl-groups, $C_{20}H_{20}NO_2(O\ Bz)_4$. The fact may be utilised for distinguishing the alkaloid of Japanese acouste from time acoustine as described on page 221.

Aconitine. Napaconitine Benzoyl-aconine

 $C_{23}H_{45}NO_{12}$, or $C_{26}H_{87}NO_7(OH)_8OCO.C_6H_5$.

Acontine is the crystalline alkaloid of the root of Acontinum Napelius, Monk's-hood or Wolf's-bane (French, Coque-luchon, German, Essenhut, Sturnhut) It exists in combination with acontine acid, C_hH_sO_a (Vol I p 452).

Aconstance is extremely dufficult to obtain m'a state of punity, owing to the famility with which it is convented into an anhydrobase, and suffers hydrolysis with formation of the amorphous base at on ine, if a mineral acid be employed in its extraction.

Alder Wright found that the whole of the alkaloid could be extracted by alcohol from Japanese acouste 100 twithout the addition of any acid; and the same appears to be true of the root of other species of acouste Thus CFB on der (Plum Centrall), xxvi

One of the best methods of proparing accusting from accounts root is that of Duquesnel (Jour Pharm et Chimie, [4], viv. 94), who exhausts the material in the cold with rectified spirit to which has been added a small amount of tartain and The alcoholic solution is distilled out of contact with the air at a temperature not exceeding 60° C, and the residue diluted with its own measure of water, and filtered from the precipitated resmous and fatty matters. The acid liquid is next agitated with ether or petroloum spirit to remove colouring-matters, and then rendered alkaline with sodium bicarbonato, which precipitates the aconitine and a portion of the amorphous bases, a large portion of the latter remaining in solution. The precipitated alkaloid is extracted by agritation with other, which, on evaporation or precipitation by petroleum spirit, doposits the base in colourless rhombic tables, which sometimes appear hexagonal in consequence of the modification of the acute angles The aconitine thus obtained is contaminated by an admixture of amorphous alkaloid, which chings to it with great obstinacy, and cannot be removed simily by crystallisation , but by converting the base into the hydrochloride, or preferably the hydrobromide, recrystallising the salt, and liberating the alkaloid by sodium carbonate, a product is obtained which, when recrystalheed from other, is vory pure.

433) has applied extraction by unacidulated alcohol to the proparation of pure acontine, and the BP process is based on the same principle. R Wright (Pham Jour, [3] xx. 375) found that chloroform alone did not extract nearly all the alkaloid from acounte root. By first moistening the not with ammonia, drying it carefully, and then percolating with chloroform, T.B. Groves obtained a much better yield than with chloroform alone. John Villia me semployed anythe alcohol for extracting continua.

For the final purification of acomithe, Dunsism and Ince (four them Sec, inx. 271) employed solution of the base in cold dilute by dechlore soid, and addition of aunc chloride in quantity sufficient to, presentiate one-fifth of the alkatod present. The amorphous alkaloid was wholly precupitated, and from the filtrate price accustine was precipitated by solution carbonate, and when crystallused from ether-alcohol was obtained in large, first, thombus prisms with truncated ends, which appeared as becaused plates under the microscops Duns tan and Ince (four Chem. Sec. in. 271) attribute to the pure acontine obtained by the above method the composition C₂₀H₄NO₂₀, which differs by H₂ from the formula of Alder W right, but the method of combustion on which both formula are based as scarcely delecte enough to decade between the two, and as hydrogen-determinations have

The following is an outline of the method of preparing crystallised aconitine ultimately piscissed by J. Williams (and posthumously published by Richards and Rogers, Chemist and Druggiet, Feb 7, 1891), being an improvement on the process previously described by him (Pharm Jour , [8], xviii 238) -The acouste root is coasely ground and macetated in the cold for three or four days with amylic alcohol, which solvent removes both the free base and its salts. The solution is shaken with successive small quantities of water slightly acidulated with sulphure acid († finid ounce to the gallon), The last washings should retain a distinct acid resotion, and the liquor should be examined to insure complete extraction of the alkaloid. The and liquid is then shaken several times with washed other, to remove amylic alcohol and coloming-matter, and then gently warmed to dissipate the remaining other. When quite cold the colution is treated with sodium carbonate, and the precapitated alkaloid filtered off, pressed, and dried by exposure to air When dry, it is boiled for some time with other (proviously washed with water and dried by potassium carbonate), and the solution filtered hot into a basin, when nearly the whole of the alkaloid will crystallise out. The ring of uncrystallisable gummy matter which forms at the edge of the dish can be dissolved by a little cold ether, in which the crystals are only sparingly coluble.

² The uncrescence appearance of accounting is regarded by Richards and Rogers as the best and most characteristic test of the alkaloid (Chemist and Druggist, May 18 1889) Crystallisation is best effected from somewhat ditte alcohol.

notoriously a tendency to be in excess of the truth, the H₄₃ formula is quite as probable as the other

Acontine is only very sparingly soluble in cold water, requiring 736 parts at the ordinary temperatus, esconding to Jurgens, and nearly ten tenes this proportion, seconding to Jurgens, and really ten tenes this proportion, seconding to J C U min o y In hot water it dissolves more freely, and is soluble in 24 parts of rectified sparis, readily in chieforem and beause, and moderately in ether; but it shaces insoluble in carbon dissulphide and percontenum spuris, and is precapitated by the latter from its solution between or either. It is not extracted from its acadulated solutions by any of these colvents

Acoustine has a slightly butter taste, the intensity of which is said to be inversely as its purity. It is extremely posisonous Solutions, sufficiently dilute to be safely employed, cause a characteristic tingling and numbness of the lips, tongue, and pharynx 1

Pure acoustine is stated by Dunstan and Inco to moit at 188° 5 C (corrected), but Duques not given 140°, Alor Wright 188°-184°, and Jurgens 1179° The material of the earlier observers was probably seasibly impure, but the went of concordance may be due in part to the mode of heating the alkinoid. Thus when slowly hested soontime melts at a lower temperature than when hested quickly Dunstan and Ince recommend the use of a bath of panellin, long enough to entirely immediate the stem of the thermometer. The bath is hested to about 150°, before the thermometer with the thing glass tube containing the alkinoid is immersed, and in kept well stirred throughout the constrain.

Acontine in the free state is dextro-rotatory, a 3 per cent solution in alcohol having a specific rotatory power of $+11^\circ$ 1 for the sodium ray. On the other hand, the salts are lawor-rotatory, the hydrochloride in aqueous solution showing $S_p = -35^\circ$ 9. Similarly

¹ Acontine is probably the most violent poson known. Ag gans it the othersy medicand does, and Ag game stat does for an adult In working with acontine, genet care must be taken to avoid the cotion of the base and its specially in the bond stats. A muster flaggement of the dust, not all to be seen, if accidentally blown into the eye, sets up the most painful initiation and lackymaton, leating some hours, while, if trabeled, although and the contraction of the dust of the desired of the dust of the desired of the dust of the desired of t

² Alder Wright states that acountine melts in a capillary table at 188°-184° (corrected). The final complete melting is preceded by a slight fritting beginning a few degrees below the melting-point, which is lowed by the presence of amorphous bases. With pure acountine very slight darkening occurs, but it is more marked with impure material.

the crystalline bydrobromide, $C_{gg}H_{gg}NO_{12}HBr+2\frac{1}{2}$ aq., gives $S_n\!=\!-30^\circ\,5$ in 2 per cent aqueous solution

SALTS OF ACOUSTING

Acontine has well-market basic properties, and forms a series of crystalinshie saits Caustie alkaline, sixed alkaline carbonates and ammonis (but not ammonium carbonate or fixed alkaline baceboates), thore down the free base from the solutions or sails as a white flocculent precipitate practically insoluble in excess of the reagent

The sales of acomtine with the mineral acids are neutral to methyl-orange and resolte acid, but may be titrated with standard caustic alkali and phenolphthalem, just as if the acid existed in a free state

Acountine Acountate exists ready-formed in acounte root. It is gummy in appearance, and crystallises with difficulty. It dissolves in water, alcohol, amylic alcohol, and chlosform, and is partially precipitated from its solution in the last menstruum by the addition of either.

Acoustine Nitrate is readily obtained by dissolving acoustine in dultae intire acid, and then adding gradually an excess of moderacily strong intro acid, when the salt separates in a bulky form, remiering the mixture semi-solud. When pressed to separate the mothal-lupon, and recrystalled from water, it forms resettes or fine showher and short premate crystals, which are colourless and transprease, but shightly efficiencement.

The acontine nitrate thus prepared has a very anomalous composition, containing as it does B₁(HNO₂)₂. The neutral astrats, BHNO₃, is obtainable as an amorphous residue by evaporating a solution in an equivalent quantity of dilute intin and

Aconstine nitiate is only sparingly soluble in cold water, but

According to J. Williams (Yaw Rod. Phon., 1888, 483), when accounts a recovered from the intrast perposed in the way it expedience in a different manner from the sugmal sileand. The extravity of the property of the propert

² A Juigens found crystallised acountine mitrate, drued at 100°, by ittrahea with caustic sikali and phenolphthaleap, to contam a proportion of mino and corresponding to the sequi-mitrate (1271 per cast), while contained of this was indicated by iosoho and (Inaugural Dissertation, Dorpat 1885)

it dissolves easily in water saturated with cubonic acid, and gradually crystallises as the gas escapes from the liquid.

Acoustine Sulphate is obtained by evaporation of its solution at a gentle heat as a viticous non-deliquescent mass, which appears under the microscope as a confused mass of crystals

Acoustine Hydroby omide, C₃₃H₁₆NO₁₂HBr, crystallises readily in monoclinic tables containing, according to Jurgens, 2½ agua.

Aconitine Hydrochloride, $C_{83}H_{46}NO_{12}$ HCl, is obtained by slow evapointion of its solution in large rhombic crystals which, according to Jurgens, contain 3 and

Acoustine Aurochlorule, CooHisNO, HAuCl., is thrown down as a vellow amorphous precipitate on adding auric chloride to a solution of aconitine hydrochloude, or to the salt of the alkaloid to which sodium chloride or hydrochloric acid has been added The preconstate is formed even in year dilute solutions, and is only very sparingly soluble in dilute hydrochloric acid. It dissolves readily in absolute alcohol, methyl alcohol, chloroform, and acetone, but less readily in other and dilute alcohol. The compound can be crystallised from alcohol, the deposition being facilitated by the cautious addition of water When pure, acoustine aurochloride melts at 135° 5 (corrected), but a very small proportion of impurity tends to reduce the melting-noint to 130°, or less From a solution of impure acoustine hydrochloride, the impurities are thrown down first, on gradual addition of auric chloride Dunstan and Ince recommend the preparation of the aurochloude and the determination of its melting-point as a reliable means of identifying aconitine, especially as the pure alkaloid can be readily recovered in a crystalline state from the compound The only successful method of effecting this, out of a large number tried, was to grind the autochloride to a fine powder with water, and add sulphuretted hydrogen water, drop by drop, tall the gold is wholly precipitated as sulphide An excess of the reagent should be avoided. The hand is then filtered, a current of air passed to remove any slight excess of sulphuretted hydrogen, sodium bicarbonate added in slight excess, and the liberated alkaloid extracted by agitation with ether

On mixing alcoholic solutions of free acomine and auric chloride, and gradually adding water, a constine gold chloride, BAUCl₂ is precipitated. When recrystallised from alcohol the compound melts at 139°.

CHEMICAL REACTIONS OF ACOUSTING

A solution of rodine in rodide of potassium produces a reddish brown or yellowish amorphous precipitate, even in very dilute (1.20,000) acidulated solutions of acontine Mayer's reagent precipitates acominas solutions, if not more dilute than 1 in 10,000, and may be used for the determination of the alksind Phophomolyhthe and also precipitates moderately dilute solutions (15000), and if the acomitine be pure the precipitate dissolves in ammonia without blue solutions are minorial without blue solutions. Phesphotungsite and behaves in ammonia without blue solutions much so with a continuous without proceedings of the solution with a continuous distriction and below 1 per cent, in strength; while platine blue inde, and potassum chromate, added, ferrocyande and ferrocyanide all to preparate acominas addutions unless very concentrated.

According to A Jurgens (Asch Pharm, [8], xxv. 127, 179, according to A Jurgens (Asch Pharm, [8], xxv. 127, 179, according a minite quantity in water scalulated with scelar and, and adding a particle of potassium nodule. On allowing the solution to evaporate, characteristic expression of some property and remain after adding water to dissolve the crystals of potassium nodule simultaneously formed.

An alcoholic solution of acomtine reduces silver nitrate, but no reduction is produced by the salts of acomtine

A mixture of solutions of potassium ferricyanide and ferric chloride is turned blue by aconitine.

Acontine, when pure, gives no marked colour-nex.tons, but as extracted from the structure and other pharmaceutroal peparations, by adding an alkalı and agristing with ether, it yields certain colour-neattons which are servicesable as supplementary tests for the acontinn-alkaloids generally (see page 242) The most characteristic property of pure acontinns is its physiological action, which may be supplemented by the reactions with sure chloride, potassum todide, and the formation of benzone acid and acontine on seponification.

As tests for the purity of acoustine, Al der Wright recommends the observation of the meltung-point, supplemented by the following —The alkaloid is dissolved in a few drops of dilute acid, pure either alded, and then access of sodium carbonate solution, the whole being well agisted in a stoppered bottle. The ethereal solution is then separated and allowed to evaporate spontaneously. When only a small volume is left, this is pound away from the deposited crystals, and allowed to evaporate completely. If the acountine were tolerably pure, the last drops of the ethereal solution will leave a crystalline readue, but if more than minute quantities of amorphous bases be present, there will accumulate in the ethereal mother-lupor, the last portions will leave a variesh or gummy residue on evaporation.

When strictly puse, accounting dissolves without colour in sulphuric

acid; and on adding a few drops of concentrated syrup no red coloration should be produced, even after standing some time

When heated for some hours to 100° C, with alcohol and caussic soda, accontine abould yield close on 20 per cent of beuzoe acel, determined as on page 234. The resulting soul should not set 120°, and should not yield any proteostebulen acel on fiscus at 250° with counts potash. This and the other reactions described on page 219 distinguish accounts from pendeaconities. Final page and the continue from pendeaconities. Final page 2014, accounts the distinguished by its crystalline form, by careful determination of the extron and hydrogen (compare page 2014), and by its behaviour with socise and benzou saliydricles In all other chancetes the two silkadost closely correspond

Acoustine is quite unchanged when heated to 100° in a vacuum, and but very slightly altered at 120°. When kept for an hour at its melting-point it loses about 10 per cent of its weight, and the resulue consists wholly of a conine, C. H., NO.,

When nomine we heated with water to 100° for many hours in a sealed two, it is hydrolysed with formation of a c on in e and be n z on c a c 14 — $C_{\rm eff} H_{\rm eff} NO_{\rm eff} + H_{\rm eff} O_{\rm eff}$. The traction is apit to be moundlets, only 85 per cent of the base being hydrolysed by heating with water maker a reflux condisenser for a few hours, the alkaloid as practically unchanged II ammons be added to the water, a small but appreciable decomposition ensues. Solutions of potassium and sodium carbonates act more powerfully, some hydrolysis courring even in the cold after prolonged standing, while on bouling nearly complete aspornification into sconne and benzosite erwise. Caustic allkalois rapidly diese the same decomposition, especially in alcobolic solution.

When acontine as heated with a ditute numeral and (especially hydrochlors early, the first scion consists in the semoval of the elements of water with formation of apo or anhydro-a contine, CAH_SNO_B. But this dehylarken is apply succeeded by hydrolysas, and formation of a conine and benzole and the state of the control of the control

ANHYDBO-AGONITINS, C₂₈H₄₈NO₁₁ is best obtained by heating acomine to 100° with a saturated solution of taitaric acid. On evaporating the ethereal solution of the base it is obtained in 214 ACONINE.

small colourless ervetals, which cohere and stack to the sides of the glass vessel in a characteristic manner. It melts at 186° 5, se. 2º lower than aconitine, and in other respects (including its poisonous properties) closely resembles the parent alkaloid. Anhydro-acontine forms crystalline salts. The aurochloride forms an amorphous precapitate which dissolves in absolute alcohol If the solution be evaporated in vacuo over calcium chloride, the compound BHAuCl, is deposited in crystals melting at 141°, but if the alcoholic solution be precipitated by gradual addition of water, the crystals deposited melt at 129°, and contain C32H48NO,,HAuCl4+H2O When this is recrystallised from dilute alcohol it is converted into a constant aurochloride, C. H. NO. HAuCl., melting at 135° 5 Anhydroaconitine gold chloride, BAuCle, is obtained by mixing alcoholic or ethereal solutions of the base and sume chloride. It melts at 147°5, and shows no tendency to pass into the aconitine salt (Dunstan and Ince, Jour Chem. Soc, lix. 284).

Commercial acoustine is hable to contain the anhydro-base, which may be removed by converting the alkaloid into the hydrobromide, and crystallising the salt from water, when the salt of anhydro-acoustine remains in the mother-liquor.

Acossins, C_aH_aNO₁, probably occurs really-founced in acontier root, and extainely in other paint of the plant. It may be obtained pure by bothing acontine with alcohole potash or soda for some hours, distilling off the alcohol, seadulating the input with hydrodhore and, and removing the beasons and by agitation with ether. On rendering the solution alkaline, and sheking with chloroform (aconine being rejudictly insoluble in eithel), the base is taken up. 1 On adding light petcoleum gradually to the chloroform aconties in the control of the control o

Aconno melts at 130°, is soluble in alcohol and chloroform, and somewhat soluble in water, but is insoluble in allythrous ether, benzene, and petroleum spirit. Both the free base and its salts resist all attempts to crystalline them. The solutions yield amouphous precipitates with the usual alkabodal teagents amouphous precipitates, which is deposited in olco-resious films on evaporating its solution in alcohol (D un at an and I n.e., Jour Chem Soc., in; 285),

¹ The author's experience is that if the alkaline liquid be shaken with other, the greater part of the basic asponification-product (aronine) is extracted, but that a small additional amount of base can be recovered by subsequent agitation with chloroform

For the solation of account from the mixed alkaloids of A Napalius, the bases are dissolved in dilute and, excess of potassium branchonate added, and the preemtated acontinne filtered off or extracted by ethics. The filtrate is slightly acubilized and precipitated by poinsion-odded of mercury, the precipitate separated, suspended in alcohol, and decomposed by subjuncted a hydrogen. On evaporating the filtered hand, the accounts is obtained as a resun which can be purified by treatment with either to service velocities and other alkaloides, solution in benzene, and precipitation by petaleum spirit. But the product is always amorphous, and predict amorphous salts

Acomic is very bitter (far mose so than econtino), but does not produce traping of the gimes, and has very little physuological scientity ($\gamma_0^{\dagger}\gamma_0^{\dagger}$ that of accountine). It is also distinguished from accountino by its unersystallivable character, its reader solubility in water and moshibility in either, and by not yielding benzoes each when boiled with alcoholic potani or sods. It reduces gold and when boiled with alcoholic potani or sods. It reduces gold and when state at the ordinary temperature and Felling's solutions on heating. It gives a blue coloration when added to mixed solutions of ferine chloride and potassium ferine/gamade

Anhydro-acoume, C₂₆H₂₉NO₁₀, is obtained by heating aconine bydrochloride to 140°. The base and salts are amorphous. It is bitter and very feebly poisonous

AMORPHOUS SAPONIFIABLE BASES OF ACONITUM NAPELLUS In addition to acoustine, the active and ervstalline alkaloid of ANapellus, and picraconitine, which appears to be occasionally present, indications of the presence of another saponifiable alkaloid have been met with by several observers. Thus Alder Wright and Luff (Jour. Chem Soc, xxxiii 318) found that the mother-liquors, from which as much crystalline acomtine as possible had been separated, contained an amorphous base showing C=66'39, and H=794 per cent, and which gave about 14 per cent of benzoic acid on saponification Wright (private communication to the author) states that it is impossible to form any idea of the pioportion of the amorphous saponitable base present, and does not regard his product as a single alkaloid, but believes it still retained aconitine, which was prevented from crystallising by the amorphous bases present. He thinks it probable that the benzoic acid produced on sanonification was mainly derived from amorphous saponifiable bases, which may possibly have been in part preexistent in the plant, but probably were chiefly alteration-products of acomtine, just as the amorphous base quincine results from the alteration of gumine.

A Juigens (Inaugural Dissertation, Dorpat, 1885) has also isolated an amorphous saponifiable base from the root of A Napsilus, and found it to contain C=6774, H=840 per cent, and to yield an unstated proportion of bemzote and and a bese allhed to acomine on saporification. BEIC, BHE, BHI, B₂H,SO₂, BHNO₂ and BHĀ were amorphous, but the very small quantity of muternal at disposal prevented any complete examination of the alkhold being made. It is probable that the amorphous asponifiable base of A Napsilus bears the same relation to acomine that quinteme beaus to quinne, and is a polymeutic of the crystalisable alkaloid. Hence the name aconicine would amprec convenient and automorpate.

J C Umney states that the anorphous saponifiable base of A Napellus produced no ill effects on him when taken in 1 giain doese.

Pseudaconitine. Feraconitine. Veratroyl-pseudaconine. CasHacNO10, or CasHacNO4(OH), O.CO.CaHa(OCH2).

Pseudacontine is the characteristic crystalline alkaloid of Acontium feron, a native of the Himalayas, and is stated to be also present in A anthona, and other species, also, according to Alder Wight in small quantity in A nanellus.

1 The remarks unde by Mr J o hn C U ma sy before the Butth Plantacetonical Ondersone of 1891 (Pharm. Jour., [3], Lut. 1926, 447; Otherset and Druggest, axxxx 293, Britaks and Ooleensk Druggest, xx 2019) contained of Jruggest, axxiv 293, Britaks and Ooleensk Druggest, xx 2019) contained of A supplies These statements, the reports of which Mr Umeny has desirated to carect, conveyed to his auditors the files impression that the recognised proportion of the macture, suponstable hase in question would settlike a decide the proportion of beamns eadit produced on asymmication, and hence would unvaluable any process of assay based on that teaction (see 283), whereas the fact is that in no invostigation, the issuffice of which have been hitherto published, has the alleged mactive loss been obtained from anountine, or in the considerable proportion errossoulty assisted by Mr Uminey, whose matrices appear to have arises in part through confinence.

⁵ According to a now recent research by Jurygens (Pharms Zarl, Sept. 1887), personutive has a constitution intermediate between accumic measurements of the polymer of a single boursyl-radead from accounts, while the dimmation of two benoply copies results in the foliantion of accounts, and the first the decomposition of accounts, on the other particular of accounts, and the property of account of the sengetive investigation spears to have been published.

3 "The raport that Morson's acontine is pseudocontine from Himalaya thin there is now deletably well desposed of, since Morson has made it known that his acontine is prepared from the tubers of collisived Aconsimon Nagallas" (H u e on a n. p. Pharm. Zech., 1884). At one time, Morson's acontine was certainly prepared from A. frow.

Pseudocountase as readily obtained pure by divadving the muture of alkahoula solated from the 100 to of A force in dhite nature, and then gradually dropping in strong intro acid with constant stirring, until, by the separation of the nitrate of pseudocountaine, the liquid becomes thick. It is then drained by means of a filter-pump, and washed slightly with water containing 8 to 10 pp. cent of intro acid. If a perfectly pure sait be required, the product is purified by re-solution in the least possible quantity of how water, colong, and dropping in stong intro acid in the sait crystallness, when it is drained, pressed, and the alkaloid liberated by treating the solution with sodium carbonic Crystallized pseudocountine contains C₂H₂N₂N₂+H₂O, but the water of anystallisation is driven off body 100.1.

Pseudocontine presents a close resemblance to acontine, both un its chemical and physiological characters? It is, however, more solublo in alcohol and either than the latter base, crystallises with I aqua, and maits without darkening at a considerable lower temporature. The melling-point is about 104*-106° Cl, but is not well maked, fritting cocurring a fow degrees lower.

When crystallised from ether, or a mixture of sthen with petroleum sputh, pseudacountine forms transparent needles and acceptable crystals, but unless the evaporation is extensely gradual the base is agit to separate as a varisals at the upper edge of sea solution, and soon forms a milk-white, cauliflower-like, crystalline efflorescence.

Pseudacontane and its salts (with the exception of the nitrate, BNO₈+3H₂O, and aurochlorade) crystaline with difficulty, and the crystalineation is impeded, or wholly prevented, by very small admixtures of amorphous alkaloid or other impurity

Pseudacontine Au ochlorada, BHAuCl., is distancily crystalline when precipitated from a dilute solution. After drying over sulphune and it can be readily crystallised from boiling alcohol in minute needles only sparingly soluble in cold alcohol and which are sullydrous when aur-drea.

Pseudaconstrue Chlosoplatsnate is soluble with moderate facility in water, and hence is not precipitated except from strong solutions. The mecuro-todide, BHgI, is amorphous, white, and very sparingly soluble.

- Anhydro-pseudaconitine and scotyl-anhydro pseudaconitine resemble the parent base in crystallising with 1H.O
- ² Pseudecontine contenns a somewhat different proportion of carbon from the other crystilline acomite bases, and the aurochlorade contains a comowhat different percentage of gold, but the best defined character of pseudacontine is its behaviour on seponification

Peendacountine is hydrolysed with great faculity. The more process of heating with dutue closel for the purpose of recrystal-lising it results in the production of a very sensible quantity of verstric a coid and pase ut do so in to (page 219). Hence only a fraction of the sikiland used crystallises out on cooling, and the mother-liquor yields venture and on addifying, adding water, and shaking with other. If freshly-preepitated psoudacountine be boiled with ammonia or acdium carbonate for a faw munutes, and the solution then endulated and shaken with either, a considerable quantity of venture and is dissolved out. When boiled under a reflux condensor for some hours with alcoholic potosh, pseudaconitine is entirely converted into versatine acid and pseudaconitine is entirely converted into versatine acid and pseudaconitine is entirely converted into versatine acid and pseudaconitine is entirely converted into versatine acid obtained approximates elected to the through elecomposition of the base. The proportion of venture and obtained approximates elected to the throughest mount (264 by pre-cent).

By heating pseudacomtine to 100° for some hours with a strong solution of tartaric and, it is completely converted into a n h y d rops = u d a c o n i t n e, $C_{90}\Pi_{e,7}NO_{11}$ (page 205), a base which closely resambles the parent alkaloid.

VERMATRIO ÁORO, GAH₂₀O₄ or C₂H₃(COH₂)₂COOH. This body has the constitution of a d in we thy 1-proto cate of true a sid H melts at 174"-175°, and can be sublimed, but is not volatile with seam. It dissolves in 2100 parts of cold water, and in 160 parts at the bothing-point, and cryst-likess from a concentrated solution at about 50° in anhydrous necelles, while crystals contening I agua rea deposited from very fultute solutions at any lower temperature. Veratrae and dissolves in alcolot and other, and is readily extincted

Possibly psoudaconitine is not the only base contained in A ferom which yields dimethyl-protocatechnic acid on saponincation. Wilght and Luff (Jour Chem Soc., xxxui 174), when preparing pure pseudaconitine nitiate by adding excess of nitiic acid to the solution of the citide salt, obtained a nitrio and mother-liquor from which no crystals could be obtained After dilution with water and separation of the precipitated resinous metter, sodium carbonate formed a copious precipitate which was freely soluble in other, but which could not be made to orystallise or yield a crystalline salt, The base was recovered from ether as a varnish, which on seponification yielded about 19 per cent of dimethyl-protocatechnic acei, and was not destitute of physiological potency, though it produced far less hp-tingling than pseudaconitine, which can be leadily obtained pure by taking advantage of its very slight solubility in a liquid containing 8 to 10 per cent of mitric acid. Since pure pseudaconitine yields 263 per cent, of veratic acid on superification, Alder Wright is of opinion that this amorphous alkaloid probably consisted of about three-fourths of pseudacourtine and other saponifiable bases (possibly alteration-products of pseudaconitine), and onefourth of non-saponifiable bases, the amorphous bases preventing the crystallisation of whatever pseudacomtine was actually present,

by the latter solvent from its accidiated aqueous solution. It produces no colonion with force chilorole. When exactly neutralised with ammonia it gives a characteristic goldanous $n+1 \vee r = n+1 \vee r = n+1$

The formation of proteoatechine and by fusion with caustic alkali forms a convenient test for pseudacontine. It is only necessary to fuse the alkalord with caustic potash and a little water at about 250° in a silver speed, acidilate the solution of the melt, extract with eitin, and test the ethercal readice with ferric children.

Other reactions of pseudocontino dependent on the vontroping group are the following—If a small quantity of the alkeloid and a fow drops of funing mitre and be evaposated to dryness, a yellow residue is ablusted, which gives a beautiful purple, and colour when movisened with a solution of causate potash in absolute alcohol. If pseudocontinue be heated with concentral sulphure and to 100°, and a drop or two of a solution of vanadum sulphate added, a violet-sed colontion is produced.

Pauronomyra, C_wH_{al}NO_p is contained in the equeous liquid obtained by spontfying pseudocontains with alcoholic potasia, acidialsting, and extincting the venture and by agistation with either 15 may be recovered by concentrating the solution, readering it alkaline by solume orbonate, and agistaing with either The base being moderately soluble in water, solume calculaproduces no precipitate in dilute solutions, and under these orcumstances either extract the base very imperfectly, but isomores certain bys-products, and on sub-sequently concentrating the allaline liquid the pseudocounce separates as a restoner mass. The last portions are readily obtained by evaporating the solution to dryness, and treating the isotatic with either, while any accuraant colouring-matters soluble in chloroform with be left unlassed void.

Pseudacomne is left as a transparent resitious varinsh on evaporation of its alcoholic or ethereal solution. On standing a few days the film from ether becomes changed into a mass of crystalline needles, but this effect is prevented by the presence of small quantities of ether, alcohol, or other foreign metters If the residue left on evaporating the etheneal solution be mostened with water a pointon of the alkaloid dissolves, while the remainder becomes opaque, white and brittle, reachly breaking up into particles having a pseudocrystelline appearance. The formation of this appaiently crystelline product seems to be peculiar to pseudosomic and lycostomics.

Pseudoconne dissolves in wates to form a solution which is strongly alkaline and very latter, but it produces no tangling of the skin or lips, and its poisonous properties are very feebly marked. The aqueous solution precipitates aliver natrate, the precipitate being reduced on heating. It also reduces ammonio-nitate of aliver on boiling, but it differs from scomme and papeocine in not reducing hot Pehling's solution, and by its solutivity in either None of the salts of pseudoconine have been obtained in a crystalline state.

Japaconitine. Sesquianhydro-japacomtine

$$\begin{array}{c} C_{00}H_{88}N_{9}O_{21}; \text{ or } O \begin{cases} C_{20}H_{29}NO_{7} & \begin{cases} O_{.CO}C_{6}H_{5} \\ O_{.CO}C_{6}H_{5} \\ C_{20}H_{29}NO_{7} & \begin{cases} O_{.CO}C_{6}H_{5} \\ O_{.CO}C_{6}H_{5} \\ O \\ O \end{cases} \end{cases}$$

Thus base is the crystalline alkaloid of Japanese acounte rock- it was first isolated by Paul and K. Ing set t (Pau-Book Pharm., 1877, 469), who ascribed to it the formula $C_0H_{\infty}N_{\odot}ok$. Lun bbe belaves it to be identical with acountine from $C_0H_{\infty}N_{\odot}ok$. The formula $C_0H_{\infty}N_{\odot}ok$ and to have the formula $C_0H_{\infty}N_{\odot}ok$. The formula $C_0H_{\infty}N_{\odot}ok$ is due to Wiight and Luff (Jour Ohem Soc., except the substitution of Paul Soc., and the sensity aspointed with production of benzone and. As the alkaloid can be extracted from Japanese acomite root by alcohol alone, without the use of and of any lund, it seems octain that the base has really the cumous constitution attributed to it, or else that the hypothetical parenthese of the formula $C_0H_{\infty}N_{\odot}ok$ or $C_0H_{\infty}N_{\odot}ok$ ($O_0H_{\infty}O_0$ suffers dehydration by the meter process of concentrating its alcoholic solution

Japaconitine is readily obtained in long rhombic crystals, and

At least two distinct openess of acousts are to be met with in the Japanese markets. Much of the root imported to Regland a sud to have been strategied in salt and runegar, and then dised in wood-sales and the sum, to protect plant of the sum of the sum, to protect plant of the sum of the sum

forms a crystallashle nitrate, bydrocliorade and hydrobronude. These salts are readily obtained crystalline by adding the dilute and to a powdered crystal of the alkaloid contained in a watch-gless, and stirring the mixture. Solidine to a clear final at first takes piace, and on further stirring a crystalline magma is formed, just as occurs with acomitine. Japacomitine is dibane, the salts containing two molecules of each

Japacontain presents the closest resemblance to acontine, both in its physical and chemical characters. Its melting-point, 184*-186*, differs only by a few degrees from that of acontine The proportions of carbon and hydrogen (compare page 205), and the percentage of gold in the surrochicald, are somewhat more tangible distinctions, but not of a very practical character. The crystalline form, as observed under the microscope, is a distinction of value, accontinue appearing in the form of hexagonal plates, and japacontinue in long columnar crystals (see illustiations to a paper WR n h a r of a and R o g e r a. Chemist and D monst. May 18, 1889)

A method of distinguishing papaconitine from aconitine, and even of estimating the proportions of the two bases in a mixture, might be based on the behaviour of the alkaloids with benzoic anhydride. According to Alder Wright, when acconstance is heated to 100° for eight hours, with twice its weight of benzoic anhydiade, it is converted into dibenzoyl-anhydroa conine, CasHarNOs(C7H5O2)2, whereas japaconitine, when similarly treated, yields a tetra-benzoylated derivative, ConHambor (CoHaO2)4 On adding a minimum of alcohol to the product, and then agitating with aqueous tartaric acid and a large volume of ether, the excess of benzoic anhydride with benzoic acid and certain impurities are dissolved by the other, while the separated aqueous liquid, when rendered alkaline, yields to ether the benzoylated alkaloids, which can be weighed after evaporating the solvent. On saponifying this product with alcoholic potash (page 204), the aconitine derivative will yield 33 40 per cent of benzoic acid, while the benzoylated japaconitine will give 50 78 per cent of the same body

Japacontine forms no anhydro-base when heated with aqueous tertaric acid

Jaraconine, $C_{20}H_{11}NO_{10}$, closely resembles aconine (page 214), and can only be distinguished therefrom by elementary analysis

Picraconitine. Cst H45NO10

This base was isolated by T. B Groves, together with aconitine, from a parcel of German roots purchased in 1874 as those of A. Napellus; but it appears doubtful whether there was not a large admixtance of some other species, or whether the roots were not of abnounced hatteres from some peculiarity of soil or claims. It is has never been met with again, unless, as is not improbable, bit better alkaoid of A gonizudatus consists of picaconstant of the contract of the contract of the possible piesence of picacontine in acounte root on must not be agnored; for, while the alkador resembles acounter too the produce of the contract of the latter base, and is piactically early physiologically, half-gram does having been taken internally without the production of any narked symptomy without the production of any narked symptomy.

Penecontine is a bittel, amorphous resul, not fusible at 100.
The dilute solutions of its salts are not precipitated by ammonia, or causise or carbonated fixed alkaloids, except on the application of heat, when the alkaloid separates as a thick congulum fusible in bothne water. Penecontine is soluble in either and chloroform

Priozoontune forms crystaliasable salts. The hydrochlorade crystaliases readily from hot solutions in fine needles. A moderately stong solution of proteomine hydrochlorade, if satarated with ammonium oblorade, becomes turbid on warming from a precipitate of the silkalontal sit, which, on continuing the heat, is wholly deposited in fine needles. The test is sine applicable to the nitrate, and probably to other salts of the alkalont.

Preasonatine gives no colout-reactions with the usual reagents. Its solutions are precipitated by tannin and Mayor's solution. The other optication is readily soluble, and the aurochloride forms a canary-yellow precipitate, not perceptibly crystalline, and exceedingly spanning soluble in water.

When boiled with alcoholic potash, picknouthne is saponified with formation of ben zoic acid and picknown of $C_{ik}H_{ik}NO_{jk}$ an amorphous base nearly insoluble in ether, forming amorphous salts, and otherwise piesenting the closest resemblance to aconine (compass footnote on page 215).

Lyaconitine and Myoctonine.

The root of Aconstan Igocolomum, a species of aconstre growing in the Alps and Humalayas, beauing yellow flowers, has been found to constant two alkaloids which differ from the bases included from other scounces So int, the products of the decomposition of these bases by alkales have not been fully studied, and some obscurity resist on other of their chanacters.

For the extraction of the bases of A. lycoctonum, Dragendorff and Spohn (Pharm Jour., [3], xv 104) exhaust the roots with alcohol acadulated with tataum and. The tancture is concentrated, mixed with water, filtered, and repeatedly arritated

with ather while still acid. The other removed traces of an acid resembling motocatechine acid, but no benzon acid could be detected. The liquor separated from the other was treated with solution beaubonate and extracted with other, which removed ly a contin to (13 per cent). Subsequent agatation with chloroform removed the remander of the lyacontine, together with myoctonine (078 per cent). The successive treatment with either and chloroform removed all but traces of alkaloid from the solution. Nather base could be obtained crystalhised.

L'accourres $^{\circ}$ was obsumed, after further purification by either of the base standed as shove, as a pale yellor resonau substance, violding a white powder, and completely soluble in ditties acids After drying m vecue, the base begins to melt at lit? 7, and is completely resonate at 114.8 (corrected), with partial decomposition. It is spaningly soluble in water, very readily in absolute alcohol, chlordorm, cubon the shiphide and betzene, less readily in either, and practically insoluble in petroleum synit. A 10 per cent solution of the base in alcohol shows a dextro-rotation, $8_{\infty} + 31^{\circ}$ 5. An aqueous solution of the initiate shows $8_{\infty} + 10^{\circ}$ 4.

The formula ascribed to lyaconitine by Dragendorif and Spohn

18 C27 H84 N2O3 + 2H2O

None of the salts of lyacontine have been obtained crystallised, the nitrate can be obtained and purified by dissolving the base in ether, and cautiously adding nitro and mixed with ether. The nitrate is precipitated, the first fraction carrying down any colouringmatter contained in the solution.

With strong sulphune and, Pacontime gives a reddsh brown colomiton, and with syrupy phosphore said a violet colomiton on warming. When treated with a mixture of 8 cc of water, 6 of strong sulphure and, and 0.3 of sodium scleenate, lyacomine is coloured a rose or pale reddish violet—a tenction which is not exhibited by the bases from other sponce of acounts.

Lyacomtine is moompletely precipitated by caustic potesh, alkaline aerbonates and ammons. Storag caustic alkalines partally decompose it Thus, when warmed for a few minutes to a temperature of 35° C with a 4 per cent, solution of caustic soda, lyacomtine dissolves, and crystalline 1 ya c o nine separate from the icquid, and may be extracted by ethen. By aquitano with chloroform a second base can be extracted, while 1 ye o c to nic sort and are resulted as a constant of the contracted of the co

¹ Also called lycaconitme

² Lyaconitine and its salts being amorphous, their composition cannot be consulered well-established. The formula attributed to lyaconino is remark-

Lyaconins, $C_{22}H_{cr}N_{C_2}+1\frac{1}{2}$ aqua, is apparently identical with the base described by H ub s ch m an in under the name of theoretonical thresholds of the control of the con

The aurochloride, platinochloride and nitrate of the base have been prepared

ACOLITITINE, a base described by H ii b s c h m a n n, is probably

able in containing an uneven number of atoms of hydrogen. Correcting it to contain H_{60} and attributing to lyaconitine the formula $C_{gr}H_{10}N_{1}O_{6}$, the puncipal reaction occurring by its reaction with soda would be —

$$2C_{27}H_{00}N_2O_6 + 2H_2O = C_{27}H_{45}N_2O_7 + C_{22}H_{16}N_4O_7$$

A specimen of "lycoctonine," from A lycoctonum, presented by Hubechmann to Fluckiger, is described by the latter chemist (Year-Book Pharm , 1870, page 99, from Archiv. der Pharm , exci) as being crystalhaed in perfectly white and distinct prisms and needles, inciting at 98°-104° without darkening, and forming a transperent glassy mass on cooling. On contact with water this mass at once crystallised. The base was soluble in alcohol, ether, chloroform, amylic alcohol, petroleum spirit and carbon disulphide By rapid evaporation from these solvents, the alkaloid formed a varnish which crystallised on contact with water, but by slow evaporation crystalline tufts were obtained. The aqueous solution of the base had an alkalino reaction and intensely litter taste. The physiological effects of lycoctonine were found to differ from those of the other acouste bases both in degree and kind As a poison, lycoctonine was found much less onergetic than acomitme. Mescurie chloride, platime chloride, phosphomolybdic acid and modide of potassium produced no precipitate in solutions of lycoctonine salts. but the base was thrown down by tannin, iodised potassium iodide, brominewater (which gave a precipitate of imorescopic needles) and the double iodides of potassium with mercury, bismuth and cadmium Potasaum morcuro-iodide threw down a precipitate which orystallised on standing. In solutions of 1 in 8000 no immediate effect was produced, but in about fifteen minutes beautiful crystals made then appearance; and in a dilution of I in 20,000 they were formed in twenty four homes The precipitate was readily soluble in alcohol. and caystallised very beautifully from the solution Meacure bromide of potassium does not affect lycoctonine solutions unless very concentrated, but both it and the mercure iedide throw down amorphous prompitates from solu tione of acondene, and do not affect narcaine colutions With potassio-iodide of bismuth lycoctomine formed a precipitate in a dilution of 1 in 40,000 Sulphure, nitric and phosphoric ands produced no colour-reactions. The nitrate of lycoctonine crystallised in tables, the sulphate in prisms Solutions of the calte were not precupitated by cauctic or carbonated alkalies, though the base itself was not notably soluble in alkalies.

sidentical with the second base extracted by Dragen dorff and Spohr from the product of the action of caustic alkalo lyncomition. It is probably a product of the further action of the alkalo on lyncomic (lyncotome). It is described as a winter powder, soluble in water, alcohol and chloroform, but insoluble in the life of the soluble of the life of the life of the life of the life of the precipitate with ammonium inolyblate. Acolytems product physiological effects similar to those of myochonies, but less reworful.

LYCOCTONIO ACID, C₂₇H₁₂N₂O₂, produced by the action of alkables on lyaconitine (or by heating the base with water or dilute and in a sealed tube), is crystallasable, and melts at 146⁻¹-148° It is sparnigly soluble in water, moderately in ether, and readily in alcohol and chloriform.

MYCCONINA, according to Dragendorff and Spoin, has the formula $C_{\rm RL}^{\rm R}N_{\rm O}$, 4-Biq. while En being taggais at as $C_{\rm RL}^{\rm R}N_{\rm O}$, 4-Biq. while En being taggais at as $C_{\rm RL}^{\rm R}N_{\rm O}$, 4-Biq. bit where being lost on drying in a current of an at 60° It is amorphous, has a latter but not pangent or impling taste, media at 145^-144°, and is dextro-totatory (8, for the alkaloid in 10 per cent solution in alcohol = +39° S, of the intrate in equecues solution 12° 2) It is difficultly soluble in water, but very soluble in alcohol, anylic alcohol, acetic ether, chloroform, henzene, and calo not desighted. Either and petroleum spirit only dissolve traces of it. The salts refuse to crystallase. Mycotonne is preepitated by most of the general reagents for alkaloids in solutions not too dhitts, and may be tituted by Mayer's solution (1 c = 0 old 176 of alkaloid).

An aqueous solution of myoctonine hydrochloride gives with excess of bromine-water an amorphous, very sparingly soluble precipitate, said to contain O₄, H₅₄Bi₆N₉O₇₉

If a fragment of myoctonine be mossened with fuming nitric acid and dired, the residue acquires a reddish brown colour on adding a drop of alcoholic potash (compare ataopure)

On heating to 100° with a 4 per cent solution of soda, myoctomue is stated by Drag-endorff and Spohn to beausimilarly to lyacontine, yielding lyocotonic acid, lyaconine, a base resembling acolyctine, and a fourth product of indefinite nature. The behaviour of myoctonine with causite alkah has also been studied by F. Einberg (Insugural Dissentation, Dorpat, 1887). When myoctonine was heated on the water-bath with 4 per cent causate soda solution, a sparmally soluble beam decomposition-product separated in crystals, which, when hitered off and purified, amounted to 24 per cent of Vol. III FART II. the myoctonuc taken. The filtents was browned, and had a peculiar pungent smell When acadulated and shaken with ether, a body exhibiting a blue finorescence was extacted, and on evaporation 3045 of a brownesh semi-crystalline residue was obtained, in which Emberg recognised bora for a call as the main constituent. The each liquid, when rendered alkaline with sodium earbonate, yielded 118 4pc erent to deliver and an additional 839 per cent to chioroform, both selvents leaving amorphous vellowish lower renduces on evaporation.

According to S al in o n ow ut z, myochonine is a powerful poison resembling curren in its achon, and acting most energetically when introduced directly into the circulation. The subestianceus injection of 0.075 gramme of the initiate produced distinct focus symptoms in eats, and the injection of 1.010 gramme always caused doubli in about half an hour. Mice were killed in three numtees by a dose of 0.010 gramme.

Atisine. C46H74N2O6, or perhaps C22H21NO2 2

Atsume is the characteristic alkaloid of Acoustum hoterophyllum, a species of acoustic which grows in the moio temperate parts of the Himalayas. The atsume exists in the root in combination with acoustic acid.

Atisine is described as white and uncrystallisable, becoming coloured and resinous on exposure to air, and melting at 85° It

To this lows, after dyring at 20°, Embring searched the formula QuHquNO₄, and land a rotation in abrothest and the Headmann's Proceedums. It is nelled a fixed and land a rotation in abrothest along the cytallune form on contact with atom. It dissolved in about 250 graits of a size, 4 of absolute aboutlo, 3 4 of allondorm, 55 of about a contact and the contact an

² The fearmin Co₂H₁N₂O₂ was deduced by the discovers of atasine, JB in ought in 0, from an analysis of the plantum as il. I was confirmed ('0) by W as 10 v or by carbon and hydrogen determinations on the free base and by analyses, the ontime of which is no not stated, of the hydrodick, which lied to the formule O₂H₂N₂O₃O₄HI+H₂O (no) 0 the other hand, O R Alder W 1; glt is found that the formule O₄H₃N₂O₃O₄ and their with determined of carbon, hydrogen, intragen and gold in the suncohloride of the base oxided by him from a small latted of that roots (Fare Jbosé Fharm, 1, 276, 422)

A heterophylikus bears flowers which are either wholly blue, or of a dirty yellow with purple times. In the bessars of India the root is sold commonly as a popular bitter tonic, under the name of Ass or Asses you. The plant and root of A heterophylikus have been fully described and figured by W as ow is or (Pharm Jone, 18] x 201, 341, 463). The root is apparently identical with

ATTISTUR. 227

has a strong, pure, butier tasts, without any accid or burning aftertasts, and is not possonous. The alkaloul as but little solution water or dilute spirit, but readily in strong alcohol, either and bonzane. When the dischlose solution is strongly diluted with water, the greater part of the alkaloul is precipitated, and the liquid froths strongly on agritation.

According to W as α w ι εx, strong sulphune and colours strain a faint roisely, which changes to read and diely blown. Nitue and produces a brown, sulphure send a red, and petessum believante a gene accloration, with a distinct reddish roisel zero. Sh in 10 γ a n n (Phan m. Jour., [3], xxv 186) obtained with some of the alkalou prepared by W as α w res α yollowine solution in concentrated sulphune and, gradually changing to a magnificent purple-red, which is state several days, but became nome-tainly varieds on adding a drop of water. No coloration was produced by mitre or hydrochions each Phosphone and dissolved the alkaloud without colour, but on warming the solution for some number at legan to solve a yellowsh valet colour. Sulphune ach and singar pounded at first a yellowish colour, which, after a few minutes, changed to vellowing red and then to carmine-sed.

The sulphate, intrate and acetato of atisine do not appear to crystalliso, but the hydrochloride, hydrobromide and hydrochlor are crystallisable and spaningly soluble salts

Ammona precipitates atisine from the solutions of its ealts in white flocks Tamun gives a yellowish brown precipitate, and potassio-menerure coldie a white proopitate, dissolving in alcohol to a solution which leaves a distinctly crystalline mass on evaporation,

Attense Hydiscidids, BHI+H,O When the precapitate of stame merus-ordule is suspended in water, and decomposed by sulphuretted hydrogen, shunng portly scales of attense hydrochia, are diposited. These thisselve in a sufficiency of het varter, and no deposited agon on cooling? The salt discovers in 318 parts of waters at 29°, and is very starnely soluble in alcohol

Atisine Hydrochloride is a white crystalline powder, more

"wakmah" or "lakmah," the former of which is regaided by Royle as the tubes of the possenous A palmatism, a view which Shimoyama (Therm Jour., [3], xvi. 86) regards as highly improbable. In anatomical characters, wakmah and atis roots exceedly correspond, and they yield the same alkaloid

¹ When the mother-inpor is concentrated to a point at which no more crystals are deposited on cooling, it will yields a presultate with potasso-ichle of moreoury, the slobholic solution of which lowes an uncrystalisable residue on evaporation. This behaviour appears to point towards the presence of a second alkalon.

soluble in water than the hydriodide. It has a strong bitter taste, but is free from the disagreeable after-taste of the latter salt.

Assay of Aconite and its Preparations.

The analytical assay and valuation of the alkaloids and other preparations of aconite vield very unsatisfactory results, not so much from the difficulty of isolating and identifying the alkaloids present, as from the uncertainty which exists between the amount and nature of the alkaloids obtained, and the physiological activity of the preparations yielding them. The most conflicting statements have been made respecting the relative activity of the actual alkaloids, even when these have been isolated in a crystalline condition, but the evidence of later observers, especially Mandelin (Phan m Joss , [3], xvi. 781), tends to show that the experiences of the earlier experimenters were due in part to the use of preparations containing a notable proportion of amorphous and relatively mert bases, to an insufficient number of physiological experiments, and ignorance of the fact that the age, sex, and general condition of an animal, besides its individual idiosyncrasy, materially affects its susceptibility to the poison. Man, again, is evidently more sensitivo to aconitine than cats or dogs, and apparently old people are more susceptible than young (compare page 236)

As a means of judging of the quality of sconito root, E. R. Squit bo (Explaneris, i. 128) recommends that a tim alice of definite section should be chewed in the lips, and the strength and length of the tingling sensation noted. A. B. Lyon's has modified that test by employing one drop of a 10 per cent uncture of the root. For highed preparations, Squit by lineas I fland anchim of a solution of the drop in the auterior pert of the mouth, previously russed with water, and holds it there for one minute, when the mouth is empired and again russed. A tenth of a minute of a 1 in 1 fluid extract, when examined in this way, should produce a distinct accounts sonsation not amounting to tingling, but very suggestive of it, and continuing more or less for fifteen to thirty minutes.

The total alkalouds contained in seconta toot can be ascertained by processes substantially idealized with those employed in preparing acoutine. The details of manipulation to be preferred have been investigated by E. H. Farr and R. Wright. (Plann Jour, [3], xxx 1037). They recommend the exhaustion of the tool by containeous percolation. One course (or 20 grammes) of the drug, reduced to coarse powder, is mostered with spirit of 0'890 specific gravity (which is preferable to other stronger or weaker alcohol), and packed in a commel prepolate, when more of the

menstruum is gradually added, and percolation allowed to proceed slowly but continuously until 8 fluid ounces (or 160 cc) of percolate has been obtained.

The tincine of acouste thus obtained is then evaporated over hot water to a low bulk, till all the alcohol is driven off The residual liquid is allowed to cool; some water added, if necessary, to reduce the viscosity, and then treated with 15 e.c. of decinormal sulphuric acid. The hound is then filtered, the precipitate washed with acidulated water, and the filtrate shaken twice with chloroform to remove colouring-matter. The separated chloroform is shaken with acidulated water to remove adherent traces of alkaloid, the squeous hand being added to the main quantity. The alkaloidal solution is then treated with a slight excess of potassium carbonate, and the alkaloids extracted by two agitations with chloroform, using 30 to 40 c c. each time The separated chloroformic solution is washed with a little distilled water, and then evaporated or distilled over hot water, the residual alkaloids being dired at 100° C till constant in weight. The alkaloids thus obtained are almost white, and viticous in appearance. Prolonged exposure at the boiling-point of water causes a slight darkening in colour1

The following proportions of total alkalous's were obtained by Fair and Wright by the above process No I sample was a root of Japanese origin; one sample was of unknown origin, and the rest were roots of A Napellus grown in Germany. The acticative metries shown in the table was determined by evaporating a measured quantity of the tincture over hot water, and drying the residue at 100°.

SAMPLE	FROM 100 0 0	FROM 100 GRANNES OF ROOT		
	Alkaloids	Extract.	Alkaloida	
No 1 (Japanese), No 2, No 3, No 4, No 6, No 6, No 6, No 7, No 8, No 9, No 10, No 10,	873 046 086 050 063 048 045 070 088 082 056 056	3 26 2 59 2 492 4 98 3 64 3 18 8 23 1 44 3 40 3 42 2 43 3 12	584 368 523 490 504 393 500 338 456 400 446	

These results show a much better yield of total alkaloids than

¹ The foregoing process is, of course, directly applicable to commercial tinging and imment of acousts. The extract should be treated with alcohol,

was obtained from the root of A Nanellus by C. R. Alder Wright. who extracted only 07 per cent, of which 03 per cent was obtained in a distinctly crystalline form 1 From the root of Japanese aconite. Alder Wright obtained 0.25 per cent of total alkaloids, of which 0.08 was civetallised Hage i found from 0.05 to 0.40 of crystallisable alkaloid, with a total yield of 0.64 to 1 25 per cent. W. Procter found 0:46 per cent of total alkaloid in American root (A Nanellus), but only 0 20 in root of German growth. From the flowers of A namendatum E. L. Cleaver extracted 0.9 per cent, of total alkaloids (bitter, net tingling), from the leaves 0.1 per cent, and from the extract of the whole plant 03 per cent. Richards and Rogers (Chemist and Dinamet, Feb. 14, 1891) extracted 0.57 per cent of crustallised acousting from dry Japanese acouste root, 014 per cent from dry 100t of A Namellus and 0.71 per cent from fresh roots (both wild and cultivated) of the same species. These results suggest a notable loss of (mystallisable) alkaloid during the process of dryppo.

All the foragoing estimations were made by farily reliable methods, and show that the proportion of alkaleds in acoustic varies widely, being probably largely affected by the time of collection, the age of the plant, and possibly by the climital and soil. The method of extraction profoundly affects the nature as well as the amount of alkaleds obtained, any hast or employed for mineral neighborhood between the proposal possible of finding with the proposal possible of the crystalline alkaleds with formation of amorrhoods lasses.

A. B Ly on a found the mostlure of acousts took to tange from 9.2 to 11.2 per cent, and the extractive yielded to alcohol to vary from 9.3 to 19.8 per cent. The alkaland from 10 grammes of the root required from 3.7 to 10.8 c., of \$\frac{3}{2}\$ Mayer's solution for its meantathom

A striking example of the effect of the process of extraction on the character and proportion of the alkalouds obtained is afforded by the following results of O Schneider (Aschiv der Pharm, cexix, No. 5), obtained with the same sample of acouster rock:

and the liquid filtered and proceeded with like that percolated from the root. The ountment can be treated cumilarly. The leaves and other pairs of the acoustic plant can be assayed in a manner cumilar to that employed for the root.

¹A still smaller yield of alkalold was obtained by Alder Wright and Rennie from the fresh (English) herb (flowers, leaves and stalks), namely, about 0 05 per cent, calculated on the dry herb, and of this only a small fraction could be obtained ovastalksed (Year-Rook Pharm. 1880. 455).

Process Employe	đ	Character of Alkaloid	Percentage		
British Pharmacopode	(1807),		002		
Morton's, .		Light yellow powder	127		
Hirzel's,			0010		
Wittatein's,		Well formed, isolated, sta-sided	140		
Hottot and Liegois',		Crystals.	296		
Duqueanel's,		Well-developed erratals	839		

The good results obtained by Duquesnel's process were doubtless due to extraction by percolation with cold alcohol, a acidulated with tartarie acid, while all the others employ more or less heat, some with and some without subdiving and

A solution of potasso-oddle of mercury (Mayer's regent) may be employed for the volunctine determination of according to the alkalouds in acid solution. The difficulty attending the use of the process is the uncertainty of the factor to be employed when there is no knowledge of the composition of the alkaloud pre-call.

Mayor's reagant may be used for the consentation of the canonic bases. The prospitation is filtered ofl, washed, suspended in water, and decomposed by a stream of sulphunetted hydrogen. The filtered luquid is trasted with an allehance callomate, and shaken with ether or chloroform; the extracted base being recovered by evanoration in the usual way to xtracted base being recovered by evanoration in the usual way.

Where the alkaloids of aconife have been extracted and obtained in a fairly pure condition, they may be detainined by titration with standard and and methyl-coange. Operating in the manuer described on page 131, the author found that very accurate detaimantions could be made. Thus 30 millynammes of erstablised

- ¹ By thation with Mayor's reagent, Zinoffaky examined the acounts continuted at Dopt in 187? Of the portions of the plants showe ground be found the flowers always inchest and the stalks powers in alkaloid, the lowest form the flowers always inchest and the stalks powers in alkaloid, the lowest concupring an intermediate place, and continuing, when fieth, about 80 per cent of water, and from 0.167 to 0.271 per cent of alkaloid. The highest need of Julyl from A. Shore-lumine. By the sawsy, appearently by Mayor's aboutton, of entire acounts plants (including the today) cellected at Dayor alkaloid or the control plants (including the today) cellected at Dayor actions of actions from the propertions of alkaloid ranging from 0.684 to 0.232 properties of alkaloid ranging from 0.584 to 0.242 properties of alkaloi
- ² In a private communication to the author, Alder Whight states that there is some reason for supposing that the crystallisable bases are apt to be more or less attaced by this freatment, and condered uncrystallisable.

soontime was dissolved in 15 cc of (neutral) ether, 3 cc, of water containing a dup of a $\frac{1}{N}$ per cent, solution of methylorange (previously rendered sensibly pink by a minute addition of acid) added, and $\frac{2}{N}$ hydrochloric acid dropped in from an accurately divined pipette, shaking well after each addition, till a permanent red coloration of the aqueous layer was obtained. Two experments made in this manner showed 2.99 and 3.10 milligrammes of acomitine, against 30 taken, while 30 milligrammes of japacomitine (not quite pure) showed 29 8 by tirtaking.

1 c c of a acid neutralises 12 94 milligiammes of aconitine.

	,,		,,	1	0 86	,,	acomme
	,,		"	3	4 14	,,	pseudacomtin
	,,		,,		0 46	15	pseudaconine
	,,		,,	٠.	2 44		jabaconitine.
	••						

The determination of the total alkaloids of an acounte preparation is in itself of little value if any as a criterion of its activity. It is rather the first step in the process of assay, the potency of the preparation substantially depending on the results subsequently obtained.¹

¹ Whene the amount of material is sufficient it is very distrible to solidate the crystalitable shalload, and it that could be effected with an approach to quantitative accuracy, it would probably furnish the most related extended the hypicalogical activity of the substance. In practice, however, very great difficulties attend such a method of examination. In the first place, these is arrays a diags, that the maximum yould of syrable may not be obtained, such boses that the activity of the preparation will be encously unfortenmented houses that the activity of the preparation will be encously unfortenmented amount of an extended which is commonly available, or on the conveniently enhanted to examination, yadde a quantity of rotal alkaloids far too small to render any method based on crystalination in particulty evaluable.

In the manufacturing klost story, when compensatively large quantities of material are waitable, a good and supple nethol of defening at least a puttal aspantian of the crystallushed attractor, and which has the advantage of being equally apphabel to accounting, no safellow —
The other of results are supplementative and papersumine, as a fallow —
The other of results are supplementative and papersumine, as a fallow as possible to the contract of the supplementative and the state of the supplementative and the supplementative and

The only pumpile of assay hitherto proposed for the acouste alkahoda, making any attempt to diseriminate between them and estimate the activity of a mixture, is that based on suponification of the active bases A method of this kind was suggested by All or Wr. 1g h, who proved that the sepantication of acoustine, pseudacounture and japacounture occurred with a near approach to quantitative accuracy (page 204).

A method of assay bised on the seponification of the crystallisable, sikeloids of acouste, has the great advantage of dastinguishing sharply between the three principal poisonous aconate bases on the one hand, and the comparatively mactive products of their decomposition on the other As it is generally accepted that aconine has only $\frac{1}{3}$ 00 of the physiological activity of accounts, and that appaceanies and pseudocomous bean a similar relation to their respective pienet alkalouds, it may be assumed that the activity of a mixture of account alkalouds is substantially represented by the proportion of crystallisable apposition base present; 2 and, therefore, the observantation of the latter with reasonable accuracy is a considerable advance towards the solution of the problem of the assay of acontine preparations?

The salt is then drained and pressed between filter-paper, dissolved in warm water, sodium brearbonate added, the liberated alkalout a tracted with ether, the othereal solution separated and evaporated, and the residue worghed.

It is true that the latter non-possonous shalled, peranontine, is separable, but it has only been net with on one occasion (1874, see page 221), unless it is identical with the imperfectly-examined latter alkaled obbuned by E. L. Cleaver from the local of the possonium oud improteoms, the amorphous alkalesis of A. ispectosure, are suposibilely, out set of no practical uniarest Both Aller W in 19th and A. Jurgons found a small quantity of an amorphous exponsibile lakelid in A. Regellus, and J. C. Un any 5 has stated that tumpshibade organization and J. C. Un and sy has stated that tumpshibade organization in program the conclusion. But neithe Wright nor Jurgons succeeded in program to be has in quantity of an opposite comments. How far these both the characteristic control to all the latter of the control to the latter of the control that is an opposite that the quantity solded was too small to allow of complete examination. How far these little-frames before the control to the state of the sponification-process of assay is uncertain, and hence the results must be regarded as tentative, except where the method is supplied to the alkaled provincely obtained in a cyretables of the order of the control of

² Alder Wright holds etrongly that all galenical preparations of acousts and amouphous alkaloids should be abandoned, and only well orystallisod alkaloids or that salts employed

It is a grave scandal that, although the acounces difference in physiologonal potency between the crystalline allicales of the scenits and the amorphous bases associated with them, or produced by their decomposition, has been long recognised, and become generally known, and while crystalline societies can be reachly prepared, that a proparation should still be sold under the manno' "moontine" which we not crystallised, and contains a large proportion of

Alder Wright's saponification experiments were made on comparatively large quantities of the alkaloids, but to be of any practical value the method of assay must be available with a quantity of acouse bases not exceeding 50 milligrammes, and should even be applicable with halt that quantity. The author has succeeded in making very satisfactory determinations on these small quantities by the following method of operating, which may be conveniently applied either to an ether or chloroform residue, or to the liquid resulting from the titration of either of these with standard acid and methyl-orange, as already described. The icudue or solution, containing 30 to 80 milligrammes of alkaloid. is treated with 20 c c of rectified spirit (neutral to phenolphthalein) and 3 c c of a solution of caustic soda in an equal weight of water The liquid is then boiled for an hour in a flask under a reflux condenser, after which the alcohol is distilled off, and the residual hourd acadulated with hydrochloric acid. The liberated benzoie or verstage acid is extracted by agriculou with about 15 cc of ether, and the ethereal solution separated and washed with successive small quantities of water, until the washings show then freedom from mineral acid by ceasing to redden litmus. The cthereal haund is then separated and transferred to a small stoppered cylinder (25 cc capacity), about 5 cc of water faintly coloured with phenolphthalem added, and 1 normal baryta-water dropped in from a finely-divided pipette until the aqueous layer acquires a pink colour, which is not destroyed by agitation with the ethercal stratnm

From the volume of standard bayta consumed, the amount of commate each isemilary from the saponification can be calculated. One e c of \$\frac{x}{x}\$ bayta heutilates 2.44 millignammes of benzon each, or 3.64 millignammes of ventuic (dimethyl-protocatednuc) each, of alkain neutralised by equivalent weights of them are, of course, dischased; and hence no grave difference results in adculating the supominable alkalond, whether benzon or venture and has been produced by the appointionation. Thus —

1 c c of $\frac{\pi}{\omega}$ baryta represents 12 94 milligrammes of acontine saponified , , 14 14 , pseudacontine saponified , , 12 44 , japacontine saponified

In three experiments, where a weight of 30 milligenames of the particular justice base. It is a question white the vale of such in the preparation as "accountes" is not an infragement of the Sule of Foot and Drugs Act, notwithstanning that the Intel® Paracogness officially tongonesses the middinate maxture as "accountes" and describes it as "usually susception."

same sample of crystallised acomtine was saponified, the havyta solution used represented 31 6, 28 3 and 30 9 of the alkaloid. In the case of japacomtine (not quite pure) the process indicated 29 8, against 30 milligrammes taken

If desired, the stration being completed, hydrochlore and may be added, when the aromatic and will be hisparded and relussived by the other. On separating this solution and allowing it to evaporate spontaneously, the weight of the send may be separated and its melting-point observed, or the other may be separated from the aqueous hquid, and the latter aculahated, largely diluted and distilled, when a separation of the benoise and vesture scale will be effected, the fourier volatile-may with the steam and the latter remaining in the rotot. This diffusions of behaviour enables seendacontinue to be recognised and estimated in presence of acontinue and paracontine.

By the foregoing method of assaying the mixed alkaloids from the fincture of A. Napellus root, G E Sectt-Smith obtained in the author's laboratory the following results —

	A	В	С	D	E	F	G	п
Weight taken, in milligrammes,	55 G	51.7	21-9	67.0	21.0	31.5	172	23 5
Alkaloud by tetration (in terms of)	69 0	067	29 1		28 8		181	24 9
Benzoic seid,	5%	40	[84		48	4	1
=Aconitine,	27 7	20 4	١.	44.5		25 7	22	1
Percentage of saponifiable alkaloid,	50 4	30 5		51.1		81.0	50	1

If desired, the base product of the seponification can be isolated by rendering the hound alkaline with solume earboants on causate sods, and agricating with ethen or chloroform. The latter solvent sortinets a trifling further quasi-lift from the hapid which has already been treated with ether. The law experiments made in this direction in the author's laboratory gave somewhat ciratic results, probably owing to the imported extraction of the bases by immuscable solvents, and the further action of the causine alkali on them.

¹ The alkalouds from a teneture prepared from the root of M foce gave, for 7 milligramment akone — By trainine, 74 of alkaloud, calculated as pseed-countine, asponified, 14 a milligrammes of venture seed by titration, against a weight of 13 of extracted by rebe. The former result regression set for 1 pseed-countino, leaving 21 6 of unsaponifiable alkaloud. The base products of the asponification extracted by reber, followed by peluodom, from the alkalime residue, amounted to 18 5 milligrammes, and nestratiled acid equivalent to 51 4 of predictioning, 600 4 of pseudocountine.

Posoning of human bougs by pure aconitine has been of compensariesy mo occurrence, but there have been numerous cases of posoning by the roots, leaves, and galencial preparations of acounts, the greater number being the result of accedent. The root has been occasionally caten in mistake for horse-radials, which it somewhat resembles (compute page 199)

The medicinal dose of the BP tincture of acomits is from 5 to 15 numms. A. Wynter Blyth considers twice the maximum dose, or 30 minims, likely to be fatal to an adult, though the least fatal dose is usually stated at above twice this measure Fleming's tracture of acomite is from three to six times the strength of the BP preparation ² The BP teniment is eight times as strong as the timeture ³ The fatal dose of aconstine is difficult to fix, as in the few cases in which a fatal dose of the pure alkaloid has been administered the quantity taken has not been known, and in the cases of poisoning by preparations of acomite there is the greatest uncertainty as to the amount of alkaloid contained therein. Headland considers to grain of acoustine an ordinary fatal dose for an adult, and the grain of the nitrate has actually caused death. Death appears to have been caused in one hour by 0 0005 gramme of accountine (Pharm Jour, [3], xx 734) Wynter Blyth considers 002 gramme or '03 gram the minimum fatal dose for an adult, when the porson is taken by the mouth, but that if given hypodermically, 0 0015 gramme would probably kill, since the whole of the poison is then thrown on the enculation at one time, and there is no chance of its chimination by vomiting Pereira relates a case in which to grain nearly proved fatal to an elderly lady Recovery has occurred after taking 24 grains, but in this case there was violent vomiting immediately, and most dangerous

J.A. Wyn ier Blyth, in his work on Posses, states that be had collected from European Heature, of the tar years pero to 1874, agisty-seven, cases of possoning by accounts in some fairs not other. In these west two cases of mustler, seven of tsurieds, and swenty seven more or less accederal. Six of the cases west from the use of the alkaland itself, ten from the root, in two cases children as the flowers, in one case the leaves of the plant were cooked and octen by mustle, in seven the tructure was mustless for sherry, brandy, or biquetr, and the romanule were caused by the intentry, the thirment, or biquetr, and the romanule were caused by the intentry, the thirment, or

² Dr Male, of Burmugham, deed from the effects of 80 drops of Fleming's tracture, taken in ten doses of 8 drops each, in the course of four

³ Dr C. Vachell, of Cardiff, has published a case of fatal poisoning by 2 grains of cett and of accounts taken in pulls. This was the maximum dose of extract according to the British Phin microperis of 1867, but in the edition of 1885 the dose is stated at \$\frac{1}{2}\$ to I grain.

symptoms for thirty hours. In the Lamson case (Guy's Hospital Reports, 1883, page 307) the victim probably received about 2 grains.

The symptoms of acouste poroming usually begin to mainfeet themselves a few minutes after the posion is taken, and are, in some respects, quite peechia and eliunateusistic. They usually, but not invariably, commence with a tingling and numbries of the lips, tongous, guins, and thicots, accompanied with a burning sensation in the storanch. These effects are succeeded by tingling and energy as existing in valuous parts of the body, pums in the aldomen, headache, vertige, and nausen, frequently accompanied by continuity of the skin, construction in the throat, finding at the mouth, partial or entire loss of voice, imparted vision, impain the cais, and feeling of tightness in valuous parts of the body, miscular tremost, and perspiration, less of numeratin power, and great pro-station generally. Sometimes there is alternate contraction and datation of the much

The most constant symptoms of acomte posoming are difficulty in heathing, noglessive mis-clair weakines, a weak intenuitional palse, and, in most cases, vomiting, especially when the poison has been taken by the month, instead of subentaneously Death issually occurs from genope, preceded in some cases by delinium and convolations. Convolutions occurred in ten cases out of metty-four collected by Dar Tucker and Reshert,* and opathodonos happens happens

¹ In a case of postoning by acouste an euseto-should be at once given, or the stomach-painp promptly used. Stimulatis may be given with advantage Ammal clarocal, to be aften wards removed by the stomach-painp, has been recommended. Strychniue and digitalis have been used successfully as artidotes, and a solution of soluted solution of postavium has been suggested.

³ In 1881, a medical man named Lamson gave his brother-in-law, P. M. John, a pout for 1/9, parlyzade before the pelva, a doss of Morona's countines, contained in a gelatin capsale. Some tearthy or thirty muniter sites, John was streak with plann in the etensoid, when he at filst called locations. He then counted, and sufficed greet pass, complained of the skin of his face being divers, of a sense of convincious in the thoust, and of leven mable to section. He stoched violently, and vometed a small quantity of dark loves and he was with difficulty kept chosen by two men. Death occurred four heavy after administration of the peason, and the visetim was conscious almost to the lest.

These symptoms probably depend largely on the dose taken. With large doses, the heart's action is arrested before the poison has had time to materially affect the excitability of the motor nerves, and the heart once stopped, further absorption is diminished or arrested.

occasionally Death from acouste poisoning commonly ensues in from two to six hours, though there is considerable variation in this respect ¹

The post-most ten appearances from acouste possoning size by no means characteristic. They are congestion of the lungs and hver, with an impected condition of the bian and its membranes. There is more or less reduces of the stomach and intestines, which are frequently found supply. Great reduces of the stomach and intestines is commented to include the secondary of the stomach and intestines is sometimes the only shormal appearance after acoustic possoning, and this does not occur when the poison has been given by the secondary of the secondary of the secondary in the secondary is successful to the best visually contains more or less blood, and the blood throughout the body is generally find or do that the colours.

TOXICOLOGICAL DETECTION OF ACCULTS

In any case of suspected possoning by seconic or its proporations, the synuptoms presented before and after death are of the utmost importances. The posson is so violent, so readily decomposed, and so wanting in delected and characteristic clientical exections, that there is but hitch hope of detecting it in the body by chemical analysis. With care, however, this may sometimes be effected, and if the chemical nections be distinctly confirmed by a physiological test, the presence of the posson may be considered definitely proved International Conference of the posson may be considered definitely proved the scanned history of the scanned alkalonds have been recovered from the turns, the blood, and the liver, and have been desteted in the stomach several months aftic death, but the posson has been destroyed in cases where the viscons have become and remained alkaline for some time from patrificative decomposition.

In cases of supposed poisoning by acouste, the stomach and intestines should be carefully examined for portions of the leaves or other parts of the plant, which, if found, may be identified by

¹ In five cases of acouste poisoning recorded by J. W Mellet, death cassed respectively in S, 10, 16, 75, and 135 minutes, while in a sixth case it did not occur till four days after the poison was taken

In the Laussen case, arrly-four hours after death, there was great indoments and unfammation of the cartine end of the intended, which had a bilatived appearance, the muonis membrane aboving in places small, slightly reased appearance, the muonis membrane aboving in places small, slightly reased to previously grey patients. The duodednum was greatly orgented, and then sever competed patients in other parts of the small intentent be been used it as membranes was evidently competed, and it is lange used so, especially towards with lived parts of the property of the p

³ It is for this reason that the symptoms of aconite possoning are described in the text at greates length than would appear necessary in a work treating of the chomist's duties rather than those of the undersol practical reactioner. comparison of their botanical characters with those of real acouste. The fragments may be washed with a little distribed waten, and masticated with the front teeth, when the persistent tingling and numbness so characteristic of acoustic will be distinctly recognisable.

For the volation of acomte bases in cases of poisoning, the suspected matters should be finely divided and tented at the ordinary temperature with strong alcohol, which should be slightly concluded with tatarca each unless already distancely aced. The liquid is strained and evaporated to a low bulk at a temperature not excoording 60°C. The resquarte, slinken with other, soparated, and rendered alkaline with solume cultousts. The alkaleuls are then extracted by agritation with either on ether-chloroform, the solution washed by agritation with water, and evaporated at a gettle butter

The alkaloidal residue having been obtained, it should be dissolved in a few drops of water acidulated with acetic acid, and a drop of the solution placed on the tip of the tongue or made the lower hn E R Squibb recommends that the quantity to be tasted should be dissolved in about 60 drops of water, which is then held in the front part of the mouth (previously rinsed) for one minute, and then discharged Another good plan is to drop the solution on a fragment of porous biscuit, which is then chewed with the front teeth. If any acoustine or other poisonous acousts base be present it will produce, in a period of time not exceeding fifteen minutes, a marked tingling sensation of the tongue and lips (somewhat similar to the effect produced by scalding the tongue with hot tea); and, if the quantity be sufficient and the hand has reached the tonsils a distinct sensation of sore throat will be observed. These effects last for a considerable tune, and are produced in a most marked and unmistakable manner by a single drop of the BP trueture of acoustc, corresponding to i grain of the root, and probably not more than Talon grain of total alkaloids The effect is so characteristic and delicate that it constitutes by far the best test for the presence of the poison. If not produced it is practically useless to apply other tests, as, in the absence of the physiological reaction they would at least be meonclusive, but, having obtained the characteristic tingling sensation, the chemical tests often afford useful confirmation, and enable the analyst to form an opinion as to whether pure acoustine or a galerical preparation of the aconite plant was taken 1

¹ An interesting case of this kind has occurred in the author's passonal experience A man of sucudal tendencies was suddenly takon violently il at a country inn. He suffered from difficulty of respiration and inability to use his timbs, especially on one side, had violent convulsions, and died before Vol. III. PART II.

The chemical tests should be applied to single drops of the activated solution placed on microscope-shides, or, in the case of the colour-tests, to the resultue left on ovaporating a few drops at a gentle heat on the made of a potechin crucible cover (compare page 145). The reactions which may give of service as the

- 1 The formation of a crystalline intrate on adding a small drop of intric acid at the end of a glass tod (page 210)
- 2 The formation of a crystalline suiochloride on adding a drop of auric chloride (page 211)
- 3 The formation of crystals of acoustine hydrochide on adding a minute fragment of potassium iodide, and allowing the solution to evaporate (page 212)
- 4 On adding cold concentrated sulphune and to the acounter results no reaction is produced immediately, but very giadually, or more rapidly on custionally warming, a deep bronic coloration is produced, passing through various shades of include brown to violet. The reaction is not produced by upon acounting.
- 5. In presence of costam impurities, which adher tenacously, ascente base develop a well-mained observed coloration, changing to cumson, when treated with sugar and sulphune acid in the manner described under morphism. The mixture of bases extended from accurate root in the admary process of assay gives this resciton very distinctly.
- 6 Impure residues of aconte bases, when treated with syrupy phosphone and, give a violet coloration when the mixture is heated for some time on the water-bath, so as gradually to concentrate the and
- 7 Acousting yields with phosphomolybdic and (Sounenschein's reagent) a yellow procipitate, which, in the presence of impurities, dissolves in anunoma with blue colour.

When the torgue-test renders the presence of an acounte base probable, it is vary deamble to make a further physicologic experiment on a small animal. For this purpose a quantity of reading or solution at least as great as that used for the totroguetest, and proferably several times as large, is made into one or more small pills with catineal, and given to a mouse or small but by the mouth. It is distinctly prefeatable to operate in this manner rather than by hypodamm empetion, in the case of such small

meetreal seastence could be obtained. Op analysis, an alkaloidal substance was soluted flow the stomesh, which gave vextely sumilar colour-reactions to the alkaloud extracted by the same means from the BP incture of acoust the alkaloud extracted by the same means from the BP incture of acoust BP is produced a distinct triggling sensation on the tongue and they, and characteristic symptoms in a monee which had calen a position of the extract made into a rull with outside.

and sensitive animals at those which must almost necessarily to employed. If two healthy (which) must be thosen, and one fed with ordinary catment made into pith, and the other with ordinary like make the pith, and the other with ordinary pills made with the alkaloudal extinct, the symptoms may be readily compared, and several objections obviated. According to Wynter Blyth, a quantity of acounte extinct sufficient to cause distinct numbons of the lips will kill a mouse or small brud if administered in this manned. J. H. Munro (Chem. Nens, alv. 110) has described an experiment in which the possioned a symrow with 01 grain of acounter not. Death ensued within an hour The contact of the grazand were nurved with the third with he transied in the crop, and the alkaloul isolated. The extract did not respond to the taste on any chemical test, but the solution, yhen seaked up in head-cumuls, and given to a tom-tit, killed the bind in two or these hours.

ATROPINE AND ITS ALLIES. TROPEINES.2

A remarkable series of intural alkinois exist in the plants of the annily Schaucec, and have been named, according to the plants in which they have been found, by ose y a mine and by ose in e, from Hyoseyamus name (healine) and Hadines, a trop inne and bellad on nine, from Alvapa belladoma (deadly mythishold), datuine, from Dubosta myopomates, see opolanine, from Sopola guesses, min, and ring orine, from Mauricapore was nais, see Allands.

1 T. Stevenson found 7272 grain of Moison's crystallised acountine, hypoderimically impected, fatal to a mouse in oughteen minutes W or m ley found Dangement's acountine oqually joient, 7272 grain proving fatal to a monse, after violent totaling and convulsions, in fluity-two munutes

2 The author is indebted to Mr A. W Gerrard and Mr R Wright for perusal and correction of this section

*Maxma.comxx, the alkaleal of the root of Mondiagons canadas, has been investigated by F B. Altros. & Annadas, chi. 312, Bor, v.v. 2169, Jour. Soc. Chem. Ind., vm. 814, 915). The analysus best accords with the formula C.,III., No.0, but does not evalued to be possibility of C.,III., No.0, representing the true composition. As extiasted by numerous and etherications, the bases to obtained as a very deliquecoust, colonities, stitucous mass, melting at 77"-79". The subplate torms small, white deliquescent piete, and the high achievale deliquescent resellers. The outcoldrined forms golden-yellow plates or needless melting at 163"-164". Bill, P.(C.) cyrishilises from het water mysolies with similar with decomposition at 138"-194". The mecuric olderide crystallines from water or alcohol in slightly voltade.

bases are distinguished by a remarkable power of dilating the pupil, and hence are often termed the "mydriatic alkaloids," though the effect of pupil-dilation, or mydriasis, is not confined to the alkaloids of the Solamaceue

More recent investigations have reduced the number of the bases supposed to exist in the Scienceae Thus, it appears that the bases isolated from A belladoma and D stranomism were simply a instance of atropine and byosepanine in varying proportions, and that hyosepanine is converted into atropine with such facility in presence of a trace of sikeli, that it is not implobable that atropine does not always pre-exist in belladomus (see page 250). Similarly, the alkialoid described as d in b or s in a supprently identical with hoseevamine, or with a mixture of that base and hyosems.

Constitution of Atropine and its Allies.

The these best-known of the natural tropames, viz, atropin, phoseyanme and hyoseina, are all isomenic, being expressed by the founds C₁H₂₈NO₃. The associated bases belladonnine and attopaume differ from these by the elements of water, and are probably analytic-bases (negs 251). All these alkaloids are readily supominable, and traces of the products of their hydrolysis are thesefone labels to pre-exist with them in the plant, or to be produced during the process of isolation. The following table exhibits the leading properties of the natural tropeners.

Base,	Formula.	Melting-	Specific Rotation	Form	Products of Saponification by Baryta			
		·c	Kotaviou		And	Визв		
Atropine,	C17H 20 NO 8	1115	+0°to-1°9	Noedles	Tropic and	Tropine.		
Hyoscyamine,	CarH ₂₂ NO ₃	108 5	-21*	Needles or	Tropic soid	Tropine		
Hyoseine,	C ₁₇ H ₂₀ NO ₃			Colouriess syrup	Tropic acid	Pasudotropine.		
Belladonnine,	C17H21NO2			Amorphous)	Isomers of tropic and	Pseudotropine.		
Attopamiue, .	C17H21NO2	Below 60	+0*	Varnish }	atropic and	Pseudotropine		
Scopolamine,	C17H21NO4				Attopic souft	Base meiting		
Benrovl-pseudo- tropine,	C ₂₇ H ₁₉ NO ₂	49	Inactive	Radiating crystals	Benzole sold	Pseudotropine.		

pedic acid, phosphotungstic acid, and iodised petusium iodide, which last yields an oily periodide Mandragorine and its salts produce inydriasis, whether introduced into the system of uncetly applied to the system of

¹ The pre-existence of atropamine and belladounine in the plants is not absolutely established

The natural tropemes are all easily saponified by treatment uth year or alkaines. By the latter (sepsecully baryta) the hydrolysis results in the formation of tropic a cid, on an iso me rethereoff and tropine or pseudotropine, in accordance with the countage.

$$C_{17}II_{29}NO_2 + H_9O = C_9H_{10}O_3 + C_8H_{15}NO$$
.

When the hydrolysis is offocted by an acid, especially continted hydrochlore acid, the topic acid loses the elements of water, and a tropic acid, $C_0H_3O_D$, results, and at a high temperature this is more or less changed into its polymers α - and β -is at ropic acid, $C_3H_2O_D$. Such products also result from the exponification of the anhydro-bases belladonume and attopanume by harris

The preferable method of effecting the saponification of the topcanes is to beat the alkalond with saturated baryta-wate to 60° or 80° C for a few hours. Carbon daoxide is next passed through the laquid sill a drop ceases to give a pink coloration with hemolybihablem. The liquid is then filtered, and the filterit acidulated with hydrochloric acid and twice shaken with either The either is separated, and on evaponition yields the acid product of the hydrodysas, on treating the equeous layer with caustic alkali in excess and agitating with either the has to product is extincted, and may be recovered by separating and evaporating the either

Thorvo Acmo, Caff. CH(CH, OH)CO OH, has the constitution of a-ph any 1-β-hydroxyp poponic act of It crystallises from hot water in needles or elendes prame, and on the spontaneous evaporation of its agueous solution in tablets which melt at 117-118° C. Tropic and is not volatile without decomposition. It has a slightly sour tasts, itsolven in 40 parts of cold water, and is soluble in alcohol and ether. When heated with a dilute solution of polessum permangumate, tropic acid gives an odou of bitta-nluond oil, and ou further treatment, bensore and is produced.

Tropic acid has been prepared synthetically (Ber, xiii 2041)

ATROPIO ACID, C₆H₅ C(CH₂) CO OH, has the constitution of a-phenylacrylic acid. It is isomeric with cumnamic acid (Part I, page 30), from which it differs by its solubility in water

¹ Except in the case of benzoyl-pseudetropine, which yields benzoic acid on hydrolysis

(1 m 692 at 19°), its lower melturg-point, and in not being preparated by uningsonous salts from its neaths solutions. Attropressor has been prepared synthetically, and may also be obtained by heating topic and with hydrodionic send, on by the direct action of funning hydrodionic send, on the concentrated buryta-water, on attopice. It expeditions from a bendoir in tablets or monoclime pursus, which melt is 106°-107°, and from alcohol in tablets or monoclime pursus, which melt is 106°-107°, are volatile with steam, and bod with decomposition at about 207°. Altopic and is vary soluble in carbon disriplinde It is considered to be razori a card by chiomes and mixture and yields for mix and plen y lacetic acid swhen fused with constructions.

Isamopro Acin, $C_{\rm in}H_{\rm in}O_{\rm in}$ is polymente with a frop ne acid, $C_{\rm in}H_{\rm io}O_{\rm in}$ as always formed tagether with that acid and track acid when atopine is heated with hydrochloric acid. Jastropine acid is always formed in small quantity when atopine acid recrystallined from hot water, and more largely if the solution be builed for some time.

Several assume to modifications of stations and exist, this as-stations can is almost excutively formed when atoms is the heated for many house to 140° – 160° m a closed flask. It forms a small warty exptsh who he mids at 237° , are very sightly soluble managam or cold housemew-water. Several part of the conditions of the amount of the amountment of the solution of atoms and the solution of atoms and the solution of atoms and the solution of a total the amountment of the solution of transition of the amountment of t

Thorstra, C₁H₁C₂H₁, ORIN CH₈, has the constitution of a tetr s hy 4r op y 1 rd 1 ne, C₁H₈, in which two of the hydrogen atoms are replaced respectively by methyl and hydroxyethyl. It is the beaue product of this staromineation of both atorogen and hysesyamine (see page 244). Tropine crystallises from absolute their in rhombia tellates, nucleus at 61⁴-62⁸ and boiling at 229°, It is hygroscopia, and very easily soluble in water and alcohol, cenaming as an oil on evaporating these solutions. Tropine is a strong testanty base and forms salts which crystalline well BH,HPUG, forms lange, ornapc-red monoclinic pursus, sassiny soluble.

in wain water, modulo in alcohol, and melting with decomposition at $108^{-2}00^{\circ}$. IHA MCI_{i} forms large yellow plates, melting that decomposition at $210^{\circ}-213^{\circ}$. The parade is a yellow precipitate, expisalising from hot water in yellow needles. On ignition with sold-lime or causite length, tropins yields at cf in Ja in in c, water and it rop ii i do ne: —C, $M_{12}NO = CH_{11}NI_{12} + H_{12}O + CH_{11}$. When heated with funning hydrochlone and to 180°, or with gleanl accine and strong sulphinus nead, it loss the elements of water and is convivated into it to p d in c, C, $M_{11}C_{11}NI_{12} + H_{12}O + CH_{12}$. When it is convivated into it to p d in c, C, $M_{11}C_{11}NI_{12} + H_{12}O + CH_{12}$ along these boiling at 162° , smelling like comme, and interesting from its relation to analytho-e-cyamic comparer page 270)

Particornovirs, C.H., NO, is vonceré with itopine, and costlatem the hydroly-as of hyo-eine, belladomine and atropamine. It forms rhombohedral cryslals, melting at 106° and boiling at 211° to 243°. It is less hygroscopic than tropine, but very soluble in water and chloudorin, and somewhat spatingly in either B.H.F.PiCl. froms small orange-red rhombo prisms, early soluble in water. Bill Mich. If nome small cryslals melting at 193°.

By treating pseudotropine with strong hydrochloric or sulphuric acid, a base isometic with troudine has been obtained

Atropine, Datume Tropyl-tropine

C17H23NO3, or C5H7(C2H40 CO CHC6H5 CH2 OH)N CH8.

Attorne is the characteristic alkaloid of Attora belladonna or deadly nightshade, though it appears sometimes to be wholly or in great particplaced by its isomer hyoscyanium.² It

1 The melting-point of the aurochloude is almost the only marked distinction between the pseudotropine produced by the hydrolyers of hyoscine and the (possibly identical) pseudotropine described by Liebermann (Bo, xxiv 2336), as resulting from the saponification of the beacoul pseudotropine discovered by Giesel in coca leaves from Java After boiling this base with hydrochloric acid for some hours the benzoic acid formed was ovtracted with ether, and the send liquid evaporated to dryness. The hydrochloride was decomposed by oxide of silver, or excess of strong caustic soda solution added, and the base extracted with other Pseudotropine thus obtained has a strong alkulme reaction, crystallises in heautiful needles, melts at 1062-1079 boils at 240°-241°, and is easily soluble in water, alcohol, and benzeno, and is precipitated by petroleum spirit from the last solution BHCl forms hygroscore needles, the colution of which is precipitated white by morenia chloride Bil.PtCla does not crystallise till the solution is evaporated nearly to drynos, but is then difficult to rechesolvo in water, and is precipitated on adding alcohol BHAuCl, forms beautiful yellow needles, molting at 225°, and easily soluble in hot water and alcohol The picrate forms easily soluble, yellow needles

² See an interesting paper by Schutto, Pharm Jour, [3], xxii 429 (from Activ. October 30th 1891, page 492)

also occurs in the seeds of Datra stramonum or the oin -s pplo, whence its name daturned * Atropine has been propued synthetically by heating together at 100°, with dilute hydrochlores end, the topic seed and tropine resulting from the hydrichies of hyoscyamine (page 244). The three too version of hyoscyamine into stropine has also been effected (page 250), though the reverse change does not anonat to have been realised.

Pure stropuse forms tults or groups of colouries or while lustrous needles, or sateular prisms. In commence it often occurs as a crystalline or nearly amorphous yellowish product. By prolonged exposure to an it gendually sequine a yellowish or darker tunt. It melts when pure at 114° G according to Let do not longer or at 116"-115° 6 according to Sch in 14t, but the commercial calkoloid often begins to melt at about 104', and is entirely meltod at 113". At a higher temperature at hopine shows again of volatility, and, according to D rag en do rft, volatilises slightly with sitem, and even with alcohol-vapour. When dry, however, atropine does not less weath by exposure to 100° C

Atopne is ofourless, but has a disagreeable inter and amy taste it is a powerful power, producing delurum and convulent (page 361) From 0.05 to 0.2 grainine is commonly fatal, and 0.001 grainine the maximum medical does for an adult. Most smaller amounts than this produce marked mydrasses or dilation of the punjul when apouled to the eve (uage 250.

Atopine as soluble in 600 parts of cold or 35 of boiling water; or, according to other authority, in 200 parts of cold and 56 of ¹ For the preparation of stepnes from bolladona, the duel leaves should be assentate less several days in cold whore, the hyand concentrated by emporttion, iterated with sodium can bonate, and agnited with Learene. The borne column is operated and agnited with diste eshiphore and, and the bound of the several days in cold with diste eshiphore and, and the catendar with colorion, the solution is which, when movel with perincitation with colorion, the solution is which, when movel with perintention of the colorion is solved in the colorion of the colorion is what the associated alkaloids ionium in the mother-inqual I is, pethapwishes the associated alkaloids ionium in the mother-inqual I is, pethapmic says to prepare atropines from balladona and Chinoform's its which

In a purate communication to the author, A W Gerrard states this prose actorine modes at 116-116 If some of the same sample be placed in water it mults at 38-54 This result is evadently due to hydration, for the solutions, after contact with water, mults at the same temperature in a capillary table, but by exposure over stong sulphune soul the allicaid loss into units, and then again modes at 11-416 Operating on the sums speaks and the same of the allicaid loss into the contract of the same of the same

solvont for the extraction of atropine from an alkaline liquid, but other is preferable for its subsequent purification and crystallisation (A. W. Gerrard). being water. The equeous solution undergoes mad change in contact with air, becoming yellow and acquiring a dissucceable smell, but without losing its texic characte. Alropine dissolves in giyeem, and is teachly soluble in alcohol, ether (60 parts), solutionform (3 parts), anythe alcohol and benvene (42 parts), anythe alcohol and benvene (42 parts), and only slightly soluble in petroleum spirit or cathon dissiphide. The solutions are noturally matrixes, or very feelly laver-oristory.

The aqueous solution of atropine exhibits a distinct alkaline reaction to litmus, and also reddens phenolphthalcin, the latter character distinguishing atropine and its isomers from almost all other known alkaloids (nece 256)

Other reactions of atropine are described on page 254, et seg By treatment with alkahes or numeral acuts, atropine readily undergoes suponification (page 245), but is not altered by buding with strong tartance acid (compare page 206). By strong intere acid it is converted into an hyd no-atropine, finese 251).

At to some Subplate, LH-SO, prepared by neutralizing stropine with duline subplimite near and evaporating this colution to dryness at 100°, is colouriess and odomics, neutral, easily soluble in vater and alcohol, but less readily in other The commercial saft is usually family alkaline, and keeps better when so made The arrows solution should be neutral of family alkaline to thirms.

According to E Schmidt, the more hypocyamine a sample of commercial stropine sulphate contains the fine is the crystalline appearance, the pure salt occurring as granular white masses. The absence of hypocyamine is shown by the solution of the sample being ortheally mactive

Ab opine hoate and calerate are employed in ophthaline surgery. Commercial Atopane and its State should be face from yellow colour, and should not become colouned on treatment with strong sulphune and or excess of ammona. The substance should leave no appreciable reaction or gentle ignition. A drop of a solution in 1000 parts of water should have an aered and butter taste, and yield a non-thestore golden-yellow precipitate with a drop of aunochlouide, which melts under bohing water. One drop of a solution of atropine in 45,000 parts of water (or less than 2 grouns per gallon), when placed in the human cryo, should cause dilation of the puril in from forty to extry minutes.

Hyoscyamine. Datume Duboisine Chi H28NO3

This base occurs in belladonna, stramonium, and other solanaceous plants in association with atropine, with which alkaloid it

¹ Hyoscyamine occurs in the seeds, leaves, and roots of heabane and other species of Hyoscyamus, in association with hyoscine. It accompanies atropine. is stomens; indeed Laden burg (Ber, xxx 3065) holds that atropine is an optically inactive base, standing to the active hyoseyamme in the same islation as meanic and stands to lawo-tertain and At any rate, by Keeping hyoseyamme at a temperature slightly above its melting-point the optical activity gradually falls, and the product is found to consist of adoptine. Occurement of processames into atropine also occurs when its cold alcohole solution is allowed to stand after a slight addition of caustic potash or soda, or even of aminoma, but as the specific tolation of the poduct nover falls below -1°9, whereas pure atropine is wholly inactive, it is processed in the product of the product of the product of the product nover falls below -1°9, whereas pure atropine is wholly inactive, it is proposed to the product of the pro

Hyosyamine forms slender colourless needles, which sometimes radiate in groups. In its solubilities and general chemical characters it presents a close resemblance to atropine, which it also simulates in its physiological effects. The distinctions between the bases are own on many 254.

Hyoscyamune Sulphate, B₂H₂SO₄, forms small golden-yellow or yellowish white crystalline scales, of a yellowish white amorphous powder, melting at 260° and dehouseene on exposure to air

Hyoscine. C,, II, NO,

(See also page 244) Tyosense occus, together with hyosenmme, in the leaves and useds of Hyosepanus when (healthouse). The "anonphous hyosepanuse" of commerce oppears in many case-to com-state thenly of hyosense. Hyosense should be causfully difficuntated from atropuse and hyosepanuse, as its mydanica effects appear to be more rapid and powerful than those produced by the latter bases, and, taken intenally in doses of $\frac{1}{100}$ gram, it produces offset alstimat from those of atoms.

Free hyosame forms a thick synup, having a close general methogological method in which it is sentenine present to the archaeon (shally nighthado), in which it is sentenine present to the archaeon of attempts, which according to Will, is not unit-questly formed from the hyver-prime during the potences toulation. Hyper-prime also come in association with attempts in the node of Datine stransaum (thorn-upplied in a secondarion with a tempt of Sepulpairy growne and Set goods), and disnote in the not of Sepulpairy growne and Set goods, and then can be not of Secondarion are commercial hyver-prime has been proposed from one or other of the stores outces, it is labels to contain more on less of the avecanted dischaed of Secondarion are commercially properly and the store outces, it is labels to contain more on less of the avecanted dischaed of Secondarion are commercially properly and the store of the avecant dischaed of Secondarion and the Secondarion are commercially properly and the secondarion and the Secondarion and the Secondarion are considered as the Secondarion and the Secondarion and the Secondarion are considered as the Secondarion and the Secondarion and the Secondarion are commercially as the Secondarion and the Seconda

E So h m 1 dt (Phorm Zed., 1889, page 583) has obtained some indication of the formation of another alkaloid besides attorned in this teaction.

² Schutte has accombly found (Phase m Jone, [3], xxxx 429), that conversion into attorneo comes when hysecyamine is kept long in solution or in the form of aurochloride, or is repeatedly erystallised from acidinted water.

³ The calmative and sedative effects of henbane, which distanguish it in physiological action from belladonna and stramonium, are undoubtedly due to the predominating alkaloid, hyoseine resemblance to hyoscyamine and alropine, but yielding pscudotropine instead of tropine on saponification (page 247). The reverse reaction has not been realised

Other characters of hyoscine, and distinctions from hyoscyamine and atropine, are given on page 254

Hyperna Upirnohomale should occur in colounless thanhue crystals, loving 12 3 per cont of then weight when dried at 100° With Vishi's test (page 237) it should give a violet colonation. Commercial hyperne hyphobromide is hable to contain a large proportion of the corresponding salt of see polamine, and, according to E Schmidt, often essentially consists of this salt.

BHAuCl₄ crystallises in prisms, melts at 200°, and is sparingly soluble in water

Anhydro-Tropemes.

Aro-Attoperse G₀H₁NO₂ preferably called anhydro-attopus ulters from attopus by the elements of water, and hence is resumence with attoparame and helladomine. It is obtained by gradually adding attopune to fusing antire and mantamed at about 50° C. On tendening the hund alkalme and extracting with ebler, the new base is desired, and is obtained on exapostang the solvent as an oil, or pursus melting at 60°-62°, slightly soluble in water, but resulty in eliholotform. Apartiques to in updatuse or irretaing to the eye, and apparently not possonous. It yields a crystalline sulphate, spannylly soluble in cold water. The chico-polatization is crystalline, and, unlike that of atropine, only sparingly soluble in hydrochilorus acid. The aux-cellois die is amorphous, and mells at 180°. Ashlydrostropine is hydroly-sed by boiling baryta-water, fonung tropine and atropine and

Atropamine, C₁₇II₁₂NO₂ (page 244), is not a constant constituent of belladoma, and, owing to the readiness with which it undergoes change, it is hable to escape recognition. It was

¹ Sconcillating, on Scopheime, CyfigNO₂, we first found in the toot of Startogridis, and has two been related in small quantities from beliadors roce, stamounen sords, and D supportable. In one case, the mydisate roce, stamounen sords, and D supportable. In one case, the mydisate roce, stamounen sords, and D supportable. In one case, the mydisate called of the last-timed plant consisted eventually drosopations. Scophiams plants of the leaves was sessimally hypospatine. Scophiams plants of the leaves and the stamous area; if dentity, while tonated introns and it behaves as a tertiany base. By behing with buylvin it is hydrolysed with formation of a tropin card and as epicalism bayer in it is playing with the support of the su

isolated by Hesse (Annales, celx: 87) by dissolving in acetic acid the alkaloids left in the mother-liquor after the preparation . of atropine, and adding common salt to the solution until a milky turbidity was produced. On standing, the hydrochloride crystallises out, and can be obtained nine by recrystallisation from boiling water after treatment with animal charcoal. On treating the hydrochloride with dilute ammonia and ether, the atropamine dissolves, and may be obtained as a soft colourless varnish on evaporation. At 60° it forms a colourless, odourless liquid, which does not lose weight at 100°. It is only sparingly soluble in water and petioleum ether, but vory readily in alcohol, ether, chloroform and benzene The alcoholic solution is optically mactive, has a bitter taste, does not redden phenololithaleur (distinctive from atropine), but colours 1ed litmus-paper blue and neutralises acids Atropamine possesses no mydratic properties, but produces a burning sensation and inflammation when dropped into the eye, whereas apo-atropine is mactive

Attopaxime is considered by H s size to bear this stime relation to hyosents that anhydros-tengine bears to its present-bears, and it is someric with belladomme, from which it differs in neally crystall inshilty of its hydrochloride and hydrobromide, a fact which affords a ready means of separating it from the other alkalones of belladoms. If the hydrochloride or hydrobromide of attopation be moustened with a minesal scot, and warmed on exposed to sunlight, the bases is readily converted into be 11d at on in ine.

Atropamme is also transformed into belladonius by solution in cold concentrated sulphium scal, or by tho mae evaporation of the solution of its sulphate. Dilute sulphium scal also effects the conression, but a preierable plan is to best atropamme with moderately concentrated hydrochlore and to about 80°. If the solution be bouled, or it baryles—water be employed as the converting agent, the belladonium first formed undergoes hydrolysis, so that stropamme and belladonium elitimately yield the sume seponification products.

BELLADONNINS, C_MLT₂NO₂ (nage 244), is someric with anhydrontropine and attopulumo¹. It is formation from the latter substance is described above. Belladonnins forms a variati-like mass, very sparingly soluble in water, but readily in alcohol, ethen, chiloroform and benzene. The salts are amorphous B_LH₂FCI₃ and BHAUCI, are yellow pulvatuent precapitates, quite modulule in cold water. Crude belladonnins is said to contain caytropine, C_HL_MNO₃ a crystalin-bable base melting a 924 in

When belladonnine is boiled with baryta-water, or moderately

According to Ladenburg the formula of belladonnine is C_TH₃₈NO₄,
and it is converted by hydrolysis into tropic acid and oxytrolune

concentrated hydrochloric earl, it is hydrolysed with formation of past of otro pine, $G_{\rm e}\Pi_{\rm e}N0$, and two a real so of the formula $G_{\rm e}\Pi_{\rm e}N0$, and $G_{\rm e}\Pi_{\rm e}N0$ and $G_{\rm e}\Pi_{\rm e}N0$, and $G_{\rm e}\Pi_{\rm e}N0$, and $G_{\rm e}\Pi_{\rm e}N0$, but as both these bodies are annophous they appear to be isomence, and not reloutical with topic and stropic and suppose and stropic and suppose the state of 100° with faming hydrochloric acid, pseu dotropin and erystallisable a tropic a cid are formed, instead of the fotegoing annophous acids

Artificial Tropemes.

When tropine (nege 246) is treated with bemory chlorate it yould be only 0,1-to pin, 0, CLL(C,H, D)EN CEL, which is the type of a series of bodies called tropenses (La den burg.) having the constitution of eaters of top in e. The natural mydratte alkaloids belong to this clars, and alcopine has nettinally been obtained synthetically be beautiful tropic with tropic and

BRECOTI-TROFINE, C.H.-(C.H. O.H. O.H. O.H. O. S. a c.y-stallastule sub-tance which forms salls very similar to those of atropine. It is a powerful local amesthetic, and when applied to the oyes produces the dilation characteristic of the natural topicures. Berropl-pseudotropine occurs naturally in certain cose leaves from Java (usgo 287)

SALICYL-TROPINE, $\hat{C}_3H_7(C_2H_4 \circ C_7H_6 \circ C_3)NCH_9$ is obtained by evaporating to dryness a mixture of salicylic acid and atropine with dilute hydrochloric acid. It is a weak poison, devoid of action on the pupil

Homomorphis, $G_{n}H_{n}NO_{n}$ is an artificial base having the constitution of a lower homologue of atropine. It is prepared by evaporating a mixture of tropine (from the sayonification of hysogramine) and namidable sund, with dithe hydrochloric acid. Mandelo eard stelf is produced by the action of hydrochloric acid. Mandelo eard stelf is produced by the action of hydrochloric acid on any gcf al.1 in, the glucosate of almonds. It is the lower homologue of tropic acid, and has the constitution of a phen yl-glycollic acid.—

Homatropine crystallises from absolute ether in prisms which melt at 98° C. It is very deliquescent, and hence is usually obtained as a syrup. It dissolves sparingly in water, but freely in ether and chloroform.

Homatropine behaves like atropine with Gerrard's test, but with Vitali's test (page 257) it yields a yellow instead of a violet coloration. With Mayer's reagent the salts yield a white, curly precipitate, and with pieric acid a yellow precipitate soon becoming crystalline.

Homatropine resembles atropine in its general physiological effects, but is less toxic, and in small doses is a true hypnotic It dilates the pupil as powerfully as atropine, but the effect subsides far more rapidly, and hence the base has proved valuable in ophthalmic surgery

Homatropiae Sulphate crystallises in silky needles The hydrochloride is crystallisable and very soluble The chloroplatinate is deposited from concentrated solutions in fine crystals BHAuCl, is described on next page

Homat opine Hydrohomskie, C_DH_MNO_oHB;, orystallises in non-deliquescent, flat rhombic persons or pletes which form wet-Like aggregations According to the British Pharmacoposia (Additions, 1990) it is a white crystalline power or aggregation of munite prostalline posts of cold water and 133 of alcoloi¹

Detection and Determination of Tropeines.

Atropine and the allied bases present a close general resomblance, alke in their physical, physiological, and chemical characters. The following table shows the principal distinctions between them.

	Atropino	Hyoscyamine	Hyoscine		
Appearance, .	Nordies or actoular prisms	Sionder, radiating necdies or crysisi- line powder	Syrup		
Molting point, °C	114 6	105 5			
Optical activity,	Inactive or feebly here- rotatory	Sa = -21°, in alcoholic colution			
Reaction of free hase with alco- holte mercuric chloride (page 256),	Red procipitate	Yellow or red precipi- tate	White precipi		
Characters of mer- curochlorido,	Gummy precipitate	Off, solidifying to plates	Amorphous or only		
Charaoters of platinochlorido,	Not pyted from 5 per cent solutions On craperation, forms monoclinic cryetals, melting at 207	Not ppted hom b per cent solutions On evaporation, forms beautiful triclinic crystals, melting at 200°	Small octohedra soluble in water, alcohol and other alco hol		
Characters of aurochloride,	Lustreles, yellow, melts at 135*-188"	Lastrous, golden yel- low scales, multing at 100"-162	Yellow prisms, melting at 198'-		
Basic product of supenification,	Tropine, melting at 61'-62'	Tropino, molting at 61"-02"	Pseudotropme, melting at 106°		

¹ "If 2 minims of chloroform be shaken with 10 minims of a 10 per cent aqueous solution, and chlorino-water be cautiously added, the chloroform will assume a brownish colour. A 2 per cent, aqueous solution is not precipitated.

The reactions of the topenes with anne chloride from the best distinctions between them Mtopine amorbiand as this with its distinction distinction that submit its mind that solutions as an amorphous or only precipitate which gadually becomes crystalline. Under the microscope it appears in resistes and other very characteristic forms—1 melts made included the water, and is deposited from its solution in boding water acuties with hydrochlorie and in numete crystals, which are bisticles after drying, and melts at 135°–138°. Hyosegamine are only in designed to the microscope in quadratic forms—1 returns its listic when dry, and melts at 160°–162°. Hyosegamine are only of the crystallises in yellow pirans which melt at 198°–200°, and no less soluble and less bustones than the physegamine sait. Homatoquae on orbito in the six first only, but you repulse the programm of the control of the Secondomnes much light of the secondomnes are described on page 251

L at a n h ur g employs the aurochloudes to separate the tropeness from each other. The attorne salt is the most module and in factional preceptation is thrown down first, while the hysesynamic salt is the most readily solidie. The alkaloids may be recovered by decomposing the aurochloudes with eightpuretted includes, adding ammonia to the filtrate, and egitating with chloroform or other.

The foregoing properties and reactions are almost the only ones affording fairly sharp distinctions between atropine and 1 is isomers. The following teactions are (when not otherwise stated) common to the three bases, and distinguish them from other altraloids.

a Dy far the most deheate test for the tropenes is their power of producing majerizans on dilation of the pupil of the eye Dilation from the application of a solution weakes than 1 in 500 causes little monuversines to the human eye, but selutions far weaker produce the effect quite distinctly, and see powerfully, and the eye of a young eat, dog, on rabbit is to be preferred. In making such an experiment, an aqueous solution must be prepared either of the free alkelod of a tis sulplate or accetate. The solutions

by the cuttons of ditton of a solution of a mounts previously distinct with time at revious of vater. About a tenth of a grain messetured with 2 manns of nitric scal, and over apacitate to the proper scale of the solution of the solution of the solution of the substant of the solution of the solution of the solution is made about a tenth of a grain be dissolved in a little vater, and the solution he made ablation with a mannous and whaten with obligations, the separated defined from all lever on warmed with about 15 minums of a solution of \hat{x} grain of packlorde of the solution of the solution of the solution of a grain of packlorde of the solution of the solution of the solution of a grain of packlorde of the solution of the solution of the solution of a grain of packlorde of the solution of the solution of a grain of packlorde of the solution of the solution of a grain of packlorde of the solution of the solution of the solution of a grain of packlorde of the solution of should be neutral or only feebly alkalme, not stonejly contaminated even with neutral salls, and not alcoluble. A drop or two of such a solution is pinced by means of a pipette or glass rod or one of the eyes, and the sare of the pupil compared with that of the fellow-eye from time to time. E R S qu in b (Epichemers, 1955) states that distinct mydrassa is produced by a solution of 000000437 gamms of stropme sulphate in less than an hour Such an intense offect is quite peculiar to attopine and its isomers (Myoseine as even more powerfully mydiatalo,) but more or less dilation of the pupil is also produced by occanne and preparations of hemicals (commo) and digitable. Acontime has a verable effect, and moctine is said first to dilate and their to contract the pupil

b Fice atropue, as obtained by evaporating its chloroformic or scheeced soliton (after liberation of the alkaloid from one of its salks by ammona), gives a red colour with phenolphitalician. This reaction is common to byeesymmia and hycence, and is also produced by the artificial base homatropine, but is not given by any other alkaloid in common use (except, seconding to Pluge, the volatile bases, comme and meckalo). Fluckinger, who first observed the pseculiar behaviour of the topenies with pinonisthalician Jama Jam, 18, xv. 601), recommends that a miningularity of the alkaloid to be tested should be placed on phenolphitaleur paper, which is then wetted with stong alcohol. No coloustion will be produced at first, but on allowing the alcohol to evapoute, and touching the alkaloid with a drop of water, a briliant of coloration will appear. Our adding alcohol the colour is destroyed, but appears again as the spart evaporates.)

c When a solution of mesouric chloride in proof-spirit is cuntiously addied to fine stropine (as oblamed by evaporation of a chloroform solution after liberation of the alkaloid by ammonia, avoining access, a red preparties is produced A. W. Gerrard, who first described this reaction (Phaim Josu, [3], xiv 718), states that the presuprate consists of merouries oxide oxide (with a trace of merouries oxide), and expresses the reaction by the following equation—2C_HagNO_+HG_U+TK_O=HgO+2C_HTK_NO_HICI. The attorpine hydrochloride reacts with an additional quantity of merium ciliorized to form the double chloride BHC, \$HICG_N, which separates in crystalline tuffs when the input is allowed to start for a two hours. In a more reconst paper (Phaim Jour., [3], xix 898) Gerrard has modified and more processly defined the meshod of making the test as follows—0-10 gram of the fine slikeloid

¹ This behaviour is peculiar. Caustic alkalies react perfectly with phenological in alcoholic solution.

(extracted from a salt by ammonia and chloroform) is placed on a watch-glass or m a test-tube, and 20 minims of a 2 per cent, solution of merenic chloride in proof-sprint gradually added and coloration is yielded at once by attorine Hyoseyamine at first becomes yellow, then darkens a little, and finally, on heating, a well-marked red precruitate is formed. If a large excess of hyoseyamme be used, merely a vellow precipitate is formed, while with a large excess of the reagent no precipitation occurs 1 Homatropine (page 253) also yields a red precipitate under the conditions of the test, but hyosene gives neither a red nor a yellow coloration or precipitate, and hence is sharply distinguished from the other tropeines Genard found no red or yellow precipitate to be produced by strychnine, brucine, morphine, codeine, veratrine, aconitine, comne, gelsemme, cafferne, emchonine, cinchondine, quimne or aumidme; though most of these bodies gave white precipitates, which in the cases of codeme and morphine became pule yellow on heating This behaviour has been confirmed by Schweissinger (Arch. Phasm. [3], xxii 827), who also states that comme gives a white precipitate (only appearing in strong solutions and soluble on warming) and scoparine a yellow precipitate with mercuric chloride, while strychnine, caffeine, arbutine, sparteine and condurangine are stated to yield no reaction. Schweissinger suggests that the test might be made quantitative for atropine by determining the mercuric oxide precipitated, but this would only be possible in the absence of alkaloids or other substances giving precipitates of any kind with mercuric chloride. The value of Geriaid's test has also been confirmed by Flückigei (Pharm Jow, [3], xvi 601), who found cocame to give a pure white precipitate which very soon turned red

d. Gornad has also observed (Pharm Jour, [3], xv. 762) the bleeston of more upon so xide from calonol and other mercurous saits by the action of atropine. If atropine be dessolved in alcohol, and four measures of water added, the solution will immediately priceptate black morecurous coafe from a cultion of mecurous intrate free from excess of and. The 15 bet prepared by adding caustic cook, drop by drop, to a solution of mercurous nitrate until a slight permanent preopitate is produced, and then filtering.

e D Vitali has observed that if a minute quantity of solid attorine be treated with a drop of fuming intric acid, the liquid

¹ Harnack (Chem Zeit, vi. 52) disputes the identity of hypergramme and dubbasine, and states that the former gives a clear solution with Gerrard's reagent, a slight turbidity appearing on continued heating, while dubbasine gives a while turbidity immediately, and on waiming a white prepriate.

evaporated at 100°, and the residue when cool touched with a drop of a freshly-propared solution of eaustic potash in absolute alcohol, a magnificent violet coloration is produced, which slowly changes to dark red and ultimately disappears, but can be reproduced by adding more alcoholic notash. The violet reaction is almost peculiar to atropino and its isomers, and is said to be produced by 0 0001 milligramme of the alkaloid Out of some eixty alkaloids examined no others were found to give a violet coloration The coloration is not produced if aqueous potash be substituted for the alcoholic solution. Strychime gives a red, brueine a greenish, and homatropine a vellow colour when similarly treated Arnold (Arch Pharm, 1882, page 564) modifies the test by moistening the alkaloid with strong cold sulphing acid, and then adding a fragment of sodium nitrate With atropine a yellow colour is produced, which, on applying alcoholic potash, changes to roddish violet and then to pale roso Strychnine gives an orange-red colour, but homatropine behaves like atropine Alkaloids which yield strong colorations before the application of the alcoholic potash (e.g., morphine, naicetine, naiceine) render the test mappheable. Fluckiger (Pharm Jour, [3], xvi. 601), recommends that I milligramme of atropine and about the same quantity of sodium nitrato should be rubbed together with a glass rod, the end of which has been moistened with a very little concentrated sulphuric acid. A saturated solution of caustic soda in absolute alcohol is then added drop by drop, when in presence of atropine a red or violet colour will be produced. When sodium mitrite is substituted for the nitrate in the above test, an orange mixture is obtained, which, on dilution with a strong aqueous solution of caustic soda, turns in succession to ied, violet and blac

E Beckmann (Atch. Phann, [3], xxx. 481) has pointed out that venatime behaves somewhat similarly to atronic with Vitali's test, but states that with introus said or a mittie instead of nitric caul, and aqueous instead of alcoholic potash, atopine gives a reddish violot coloration, and venatime a yellow one

f When atcourse is heated to the holling-point with a mixture of equal measures of glaund acests and stong subplume acids no coloration is produced, but after a time the liquid exhibits a well-mixed yellowah or however higeon fluorescence. After cooling, the liquid has a pleasant aromatic adour in addition to that of meetic acid. The behaviour of other tropense with this test, which is due to E. Be e k m an n, does not appear to have been recorded Venature gives a similar brownish fluorescent liquid, but during the previous heating the solution acquires an intense cherry-red colors. g. According to Δ . Wy ner E lly thi, if a particle of strooms by

treated with a few drops of concentrated baryta solution, the liquid evaporated to dryness, and the residue strongly heated, an agreeable odour resembling that of hawthern-blossom will be perceived

- h According to the Germans Phas macoposa, if at least 0 001 gramme of atropine sulphate be heated in a small test-take until white vapours appear, and 1 5 gramme (=08 cs.) of sulphure and be then added, and the heating continued until the muture begins to turn brown, on then adding 2 cc of water an agreeable odour will be pnearered, and on further adding a cystal optoassum permanganata, the odour of bitter-almond oil will be obtained.
- A saturated solution of bromine in hydrobromic acid 1 gives with attornine and its salts, even in very dilute solutions (1, 10,000). a vellow amounhous mecanitate, which in a short time becomes crystalline The precipitate from somewhat strong solutions of the alkaloid disappears after a time, but is immediately reproduced on adding more of the reagent. The precipitate is insoluble in acotic acid, and only very sparingly soluble in a large excess of the mineral acids or fixed caustic alkalies. It is even produced from a solution of atropine in concentrated sulphuric and microscopic appearance of the precipitate is highly characteristic. exhibiting under a magnifying power of 75 to 125 diameters lanceolate, leaf-like erystals, grouped together like the petals of a These forms may be obtained by the spontaneous evapuration of a drop of liquid containing only washed grain of attopine, If not produced, a drop of water should be added, and evaporation repeated T G. Wormlev, who is the observer of the reaction. considers the formation of the crystals quite characteristic of atropine or hyoscyamine. Most alkaloids give vellow precipitates with Wormley's reagent, but all these deposits, except those produced by atropine, hyoscyamine and meconin, remain amorphous, and that produced by the Inst-named substance has quite a different unicroscopic appearance from those formed by the mydriatic alkaloids The behaviour of hyoscine with Wormley's reagent has not been recorded.
- j A solution of todamo jancolate of poissesum throws down, from solutions of stropues, prosegramme and hyosena, eachlated with hydrochlorre and, the whole of the alkaloud as a redshab brown or dark green amorphous presuptate of the tri-oddie, insoluble in soste ead, but somewhat affected by other acads. Dunstan and Ransom (Pharm. Josus 1 [31, xor 629) recommend the respect for
- Wormley states that m the absence of hydrobiomic acid, a solution of bromine in alcohol may be used. A solution in hydrochloric acid would appear preferable.

this purification and determination of atropine and its somers. For this purpose they dissolve the alkalod in dilute hydrochlora, and add sacess of a stong solution of rodine in potassium rollule. The precipitate at none agglomentes, and is filtered off, sight washed with the solution of rodine, and then decomposed by pouring on the filter a solution of sodium thousulpinks, only dissolves it to a colourless liquid, from which the alkalod is recovered by addition of samons and agglation with oblications.

- h Mayer's reagent precipitates atiopine and its isomers from solutions not too dilute, and has been employed with limited success for their quantitative determination. The characters of the precipitate and the best method of operating have already been fully described (lage 140 et seq).
- I Potassio-iodide of bismuth and potassio-iodide of cadmium precipitate atropine from highly dilute solutions. Their reactions with the isometic alkaloids have not been recorded.
- m Phosphomolybdic and phosphotungstic acids precipitate atropine and its isomers from somewhat dilute solutions, and are of service for concentrating the alkaloids and separating them from other organic matter.
- n An alcoholte solution of piece and yields a yellow amorphous preceptate in solutions of stropine which are not too dittie. The preceptate becomes crystalline after a time, and appears under the increascope in highly chancelevate forms. With hyosynamic, larence and yields an only preceptate, rapidly soludifying to rightangiel lamine, very similar to those formed by stropine poracity.

The reactions of atiopine and its isomers with other reacents are not characteristic. Potassium nodide, thiocyanate, ferrocyanade, ferricyanade and chromate fail to precipitate even concentrated solutions of these alkaloids

Altopine and its allies are not removed from acidinited solutions by agitation with immiscible solvents. From solutions reindered alkaline by ammonia, or an alkali-metal carbonate, they are readily and completely extracted by chloroform, and with less faculity by either. The separated solution may be evaporated, and the readine drond without loss at 100°. The bases thus soluted are distinguished from all other well-known alkaloids by their power of reidlening phenolphthalen (test b), and (with the exception of hysesine) giving a red precipitate when warmed with an alcoholic solution of meature achieved the second of the solution of meature and the solution of th

TOXICOLOGICAL DETROTION OF ATROPINE AND ITS ALLIES

Atopine, hyescyamine and hyescine are all highly poisonous cases of poisoning by the pure alkalouls are rare, but both criminal and accidental poisoning by the plants of which they are the active principles have been frequent, and, in India, poisoning by stramonium has achieved the position of a profession.

The symptoms of possoning by atropine and its isomes are thus described by A. Swame Raylo: —Heat and dripnes of the mouth and throat, nansee, vomiting, galdiness, indistinct or double vision, delirum, great exortenent and retilessuess, convulsions followed by drowniess, stupor, and lethaugy. The pupils are much dilated and the eyes meserable to light Occasionally the pupils are routracted during sleep, although dilated in the waking state. The symptoms often come on very soon after taking the posson, while recovery may be delayed for several days, or even weeks. The symptoms of possoning by stramonium are very similar to those produced by belladonns and hyosyamus, but more severe Ringing in the ears, dryness of the thinat, and flushed face are only symptoms. Delirum of a volent kind, with spectral illustors, comes on rapidly, and the pupils are widely dilated. There is often panalysis of the lower extrements.

The post-mostem unications of possoning by stropine and its isomers are not characteristic, except that the pupils are distill. The bism and its membranes are found congested. Where solid parts of a solinaneous plant have been caten the fragments of often be found in the stomach, and identified by their botameal and meroescone characters.

The detection of atropue and its isomers in cases of possoning may be effected by the Stas-Otto puccess Heating with alkales or mineral acids must be avoided, or the alkaloud may undergo hydrolysis (nego 245). Hence tatarate or acctice acid should be used to eatify the matters to be examined. Ammonia or a carbonate of alkal-metal should be used to hierate the alkaloud, and ether or (preferably) chloroform employed for its extraction. The tests most serviceable for the recognition of atropine and its isomers in cases of rossonium services.

- The dilation of the pupil (page 255)
- 2 The reactions of the free alkaloid, as obtained in the chloroform-residue, with phenolphthalem and a spirituous solution of mercuric chloride.

¹ The symptoms of attorne posening, especially in children, are not unlike theory consistency of activities and the garanton debruum and hallucinations of an adult are very similar to those of delarium temons.

- 3 The reaction of a solution of the alkaloid with bromine (page 259), and the microscopic appearance of the precipitate
- The production of a violet colour by Vitali's test (page 257).
- 5 The evolution of an agreeable odour when the alkaloid is evaporated to digness with buryta-water, and the residue heated.
 - 6 The microscopic appearance of the pierate.
- Atropine does not appear to suffer change in the body after death. It has been detected after a considerable interval of time, Ptomaines having a mydinatic action have been met with.

Belladonna, Henbane, and Stramonium.

At one belladowns or deadly inghthshada. Hyas yamus noper or henhous, and Datura stromouns or thorn-apple 2 are that chief sources of the tropennes, but these or animar sikaloids are found in a number of allied speeces, and the posenous silvant of solanne occurs in all the speece of Solanum, as well as in other members of the Solanace or the Solanace of Solanum, as well as in other members of the Solanace or the Solanace.

In addition to the alkaloids, which are probably in combination with malic acid, belladonna root contains cellulose, staich, sugar, mulin, asparagm, fatty matter, a fluorescent substance, and

- ¹ French, la Billadone, la Morelle furieure, German, Tollkirsche, Wolfskrische, Tollkiraut
 - ³ French, la Jusquiame, German, Bilscala aut
 - 3 Fronch, Stramoune, German, Stechantel
- ⁴ A minute proportion of an alkaloid, apparently identical with hyosoyamine, has been found in lettuce by T S Dymond (Proc. Chem. Soc., 1891, p. 165)
- 5 The fluorescent substance contained in belladonna root, and present also m the leaves and stalk, is called by H Kunz (Arch Pharm, [3], xxm 722) chrysatiopu acid, and is said to have the formula C12H100; H Paschkis (Arch Phon m. [3], xxm 541, xxiv 155) has isolated what is apparently the same body from the bennes of upc belladowns, and ascubes to it the formula C10H4O4 He considers it identical with the scopeletin obtained by Eykman from Scopolia supogua It forms pale vellow, rhombio prisms or needles, melting at 198°-201", and subliming without decomposition when carefully heated It disselves in about 80 parts of hot water, more spaningly in cold water and other, but readily in acetie acid, alcohol, obloroform, amylio slcoliol and benzenc. It is extracted by the last three solvents from its aqueous solution The aqueous, alcoholic and ammoniscal solutions exhibit a spicified blue fluorescence when drints, and emerald-green when concentrated. The fluorescence is destroyed by acids

 Ferric chloride gives an emerald green
 coloration changing to cobalt blue

 Folling's solution and ammonio-nitiato of silves are reduced on warming. In moderately concentrated nitite and the substance dissolves with yellow colour, changed to blood-red by ammonia (This reaction resembles that of a sculin, observed by Sonnenschein)

Kunz isolated chrysatronic acid by treating the extract of belladonna with acid and agricating with ether. On evaporating the other, and washing the

a net colourng-matter called at rosm, which is also found in considerable quantity in the besites. The proportion of statch in young belladoma roots is considerable, but it is present only to a limited extant in older and more swoody roots, and, according to a limited extant in older and more swoody roots, and, according to Morz, is almost entrally alseent during summer. The following analyses of air-dry belladoma noots are due to E. M. Hol mes—

			Wood	y Boots	Set	Roots
Moisture	, .		7 94 1	mer cont	10 28	per cent
Soluble	ash, .		3 43	.,	2.20	.,
Insolubl	e ash,		4 60	12	8 68	,,
Alcohola	o extract,		22 53	"	29 87	"
Aqueous	extract.		15 96		10 50	

Belladoma herese contam cellulese, chlorophyll, alkalodal saits, fatty and tesnous matters, & Choine is present, and, according to Bilts, asparegm sometimes crystallises from the extract after long keeping, but the crystals observed by Attified leads of potassium nitrate and chloride. By dialyses, Attifield solated potassium intrate and chloride. By dialyses, Attifield isolated from the herback and square presses of an organic sait of magnesium Kunz found 0.6 per cent of succine and in an extract prepared from the herbaccous parts of belladoma. Pinckiger found this sain of day belladoma leaves to amount to 14.5 per cent, and to consist chiefly of the cu-bonates of calcium and the alkabi-nated to

With regard to the alkaloids of belladonna, O Hesse (Annalen, cclxi, 87) states that in his experience the herb of cultivated belladonna contains atronine almost exclusively, but that it is associated with other alkaloids in the leaves of wild plants, and especially in the roots of both kinds. In an old root, Hesse found much hyoscyamino but no atropine E Schmidt (Pharm Zeit, 1889, page 583) found hyoscyamine but no atropine in full grown roots which had been kept for years. In roots of one year's growth he found both atropine and hyoscyamine, but the latter alkaloid only in fresh old roots. The leaves of wild belladonna contained much hyosevamine and a little atropine, while the tipe bernes contained atropine only E Schmidt has found both hyoscyamine and hyoscine in Scopolia atropoides and Scopolia japonica,1 and traces of an alkaloid having a mydinatic action in Solanum tuberosum, S nigrum and Lycium barbarum Mandragorine, the alkaloid of Mundragora vernalis, is myditatic and possibly isomeric with atropine (page 243)

crystolline residue with cold ethen, chrysatiopic acid remained, while 1 ou cotiopic acid, $C_{17}H_{20}O_{5}$ dissolved. The latter is a bitter substance, crystallising in microscopic pisms which mielt at 74°

³ Dunstan and Chaston found the alkaloid of Scopolia carmiolica to consist of hyose; amine with a possible trace of hyoseine.

A W Gernard (Year-Book Pharm, 1881, 1882, 1884) has published a number of valuable observations on belladonna, in which he found the following percentages of alkaloid —

Age of Plant					Wild.	Plant	Oultreated Point		
40	e of .	Plan	•		Root	Leaves	Root	Leaves	
Two years,			_		 260	481	207	320	
Three years,					881	*407	870	151	
Four years,					410	510	818	491	

These and other observations of Gerrard show that the leaf of belladoma is the part of the plant ruches in alkaloid, the root, fruit, and stem coming next in the order stated. The results of A by any and stem coming next in the order stated. The results of A by the same dustunction, for in twelve samples of (au-tired) leaves the proportion of alkaloids varied from 0.4 it to 0.69 per cent, and fifteen samples of roots from 0.47 to 1.35 per cent. The extraction matter in the leaves (au-tired, and treated with 6.6 per cent alochel) ranged from 6.6 to 1.21 per cent, and in the roots from output of the results of the roots of mostime. Lyons states that the pressed leaves do not suffer deterioration when kept for as xyaars.

R Kordes found 058, and von Gunther 083 per cent of alkaloid in beliadonna leaves, while Lefoit gives the average yield from 8 specimens at 0436 per cent

As the general result of published investigations, F arr and Wright state that the proportion of alkaloids m good specumens of commercial belladonna leaves mages from 0.30 to 0.87 per cent, their own expainments varying between 0.30 and 0.90, with an average of 0.49 per cent Gaunan leaves are distinctly poorer in alkaloid and extractive matter than those of Hogishi growth, and hence the BP direction to prepare the tincture from the leaves of "plants grown in Strian" should be strictly observed As one part of belladonna leaves produces 20 parts of the BP criticate, it follows that the proportion of alkaloid in this preparation averages 0.025 per cent, which strength might advantageously be adopted as a standard

For the assay of belladonna root, Dunstan and Ransom (Pharm. Jour, [3], xiv. 623) recommend extraction in the

¹ The influence of age on the proportion and nature of the alkaloids of belladonna has also been studied by Schutte (Pharm Jour, [3], xvn 429). following manner -20 grammes of the dry and finely-nowdered root is extracted by hot neceslation with a mixture of equal volumes of chloroform and absolute alcohol 1 If an extractionapparatus be used about 60 e.e. of the mixture will be recovered The solution is agreed with two successive quantities of distilled water, using 25 c.c each time. The separation of the aqueous liquid from the chloroform occurs promptly and completely on warming the liquid slightly. The chloroform retains nearly the whole of the colouring-matter, while the alcohol and alkaloids (as salts) pass into the water? The aqueous layer is separated, and agitated once with chloroform to remove the last traces of colouringmatter, after which it is rendered alkaline with ammonia, and agitated twice with chloroform, using 25 c.c. each time, to extract the alkaloid. The separated chloroform is agreeted once with water rendered faintly alkaline with ammonia and then evaporated. the residue being dired at 100° till constant in weight alkaloid thus isolated is obtained as a perfectly transparent fused mass. It is soluble in water, and the squeous solution gives mecipitates with Thresh's, Mayer's, and Sounenschem's reagents (pages 136, 138) It gives a faint white precipitate with mercuric chloride, and a conious white precipitate with gallottunic acid cautiously added This last precipitate is very readily soluble in a slight excess of the reagent, a distinct trace, however, invariably nemaining undissolved (Farr and Wright).3

Instead of weighing the isolated alkaloid it may be titrated with standard acid and litmus (or methyl-orange) as recommended by Gerrard (Year-Book Pharm, 1884, page 447).

Dunstan and Ransom (Year-Book Phorm, 1885, page 391) recommend continuous percolation with boiling absolute alcohol for the extraction of the alkaloids from belladonna leaves, and they proved that the leaves thus treated yielded no further

¹ Chin-nötma ilono ettracis the alikulosis vay meomplesisly Alcohlo employd, allono divolves monho ekuncture matter shali, impodes the subsequent extraction and purification of the alikulosis. If sectical spurit material of absolute alcohol be employed in admixtura with chlorofoun, the water present causes swelling of the material, and the progress of the extraction is made to the control of the extraction of the control of the extraction and alcohol excommended by them completely extracted buildings in root, and that pure stropine was not appreciably affected by prolonged boiling with the solvent.

² Although Dunstan and Ransom found the whole of the alkaloids to pass into the aqueous liquid, A B Lyons points out that it is desirable, as a mecantion, to make a small addition of sulphuric acid to the water employed.

⁹ The alkaloids from stramonum behave similarly, probably owing to the presence of a small quantity of another (third?) alkaloid quantity of alkaloul when holied with dilute hydrochlorus acid, or when mixed with lines and extracted with chloroform. Front on the extract obtained on evaporating the alcohole liquid, they found it impossible to remove the whole of the alkaloid, even by many successive textiments with water or dilute hydrochloric acid. They therefore recommend that the alcohole liquid should be diluted considerably with water acidiates with hydrochlorus acid, and the liquid then shaken repeatedly with dilotoform to remove the chlorophyll and fail. From the liquid thus purified the alkaloids can readily be obtained pure by adding excess of ammonia and extracting with chloroform.

A modification of the foregoing process is recommended by D unitain and Rainsom for the assay of the solid echact of beliadenia. Two grammes should be warmed with thinte hydrochloric and until as much as possible is theselved, when the liquid in filtered through cotton or glass wool, and the readine well waised with both dutuch hydrochion, and The filtsate is repeatedly shaken with chloroform to rainove chlorophyll, then ammonia added, and the liberated alkalonia extracted with chloroform.

The functure of balladonna can also be assayed by the foregoing and process after evaporating off the greater part of the alcolol, and the same remark applies to the final extent It is, however, many cases preferable to frest the clear liquid to access with amount and chlorotorm On subsequently treating the separated chlorotorm on the contract of t

¹ J Williams suggests that it would be better to employ ether at this stage of the process

³ Farr and Wright have shown that the strength of also his sed in section beauting the due has halted effect on the proportion of alkelond in the indicate, though it very greatly effects the proportion of maniagnosis and coloring-matries extracted, and the former of these impetch be speaked to find coformed and aqueous layers. The difficulty may be overcome by evaporation that the tructure to a strong and tracting it with strong alcohol, which proposition the muclage, and the filinate gives on evaporation a liquid which can be result's dail twit.

Farr and Wight find it impossible to sensive the whole of the alkalonds of beliadoms (and hashnas) by repeated agitation with either in present and interest of the alkalonds of aminoms, at least 20 per cent of the total remaining unortacted by other, an unattained solvent for extracting mydrasto alkalods, and the centre of the other with him to alkalods, and the centre of certard and others who have seet it as probably below the truth for fact, Gernard himself states that wreal extractions with either an encarred and results at a rich, as sub-spectrally reduced its mannounced solution mental with criter on texture and, evaporates it to a small volume, it sats it again with amongs, and again against with either which we will be a small control to the control of the control of

the chloroform, while the alkaloids can be recovered in the pure state by rendering the acid liquid again alkaline, and agritating it with chloroform

A W. Gerrard has employed substantially the same process as the above for the assay of the root and leaves of hankane (Pharm Jour, [3], xm 212, xm 213). The substance is direct at 100°, powdered, and exhausted with proof-spint. The spint is abstilled off, and the semi-fluid extract treated with water containing 1 per 1000 of hydrochloric and, filtered, and the filtered rathes diluted to 100 ce. The alkaholas are extracted by ammonia and chloroform in the usual way, punified by solution in ether, and agitated with hydrochloric and, again hierated by ammonia, extracted by ether, and determined in the alkaholad resulte by titeration with denormal hydrochloric and. The following results are recorded,—

Variety of Henhane	Part Used.	Where Grown	Yield of Alkaloids per 1000
Biennal,	Roots. First year's leaf Second year's leaf, Leaves and tops Rathe holb First year's leaves (old) Second year's tops (old)	Middlesex. Sussa, Lincolnainte Lincolnainte Lincolnainte Sussa, Middlesex Middlesex Lincolnainte Letcesterahire. Suriey Middlesex Cermany France England. England	1 1002 1 1500 1

Gerraud's experiments appear to show that a considerable falling off in the alknoids value of the leaves occurs with age the consuless that bright-coloured, well-meserved henbane, whether annual or binand, can be related no to yield good peparations, while old and dank-coloured leaves, contaming stalks and furnt, should be avoided He regards the first year's root of beneding Hypergamus raper as much incher in alkaloids than the herbacous protions of the plant, but both as much poorer than the respective parts of belladonia. * Hypergamus allius is much used in the south of Europe, but no greates extrength as attributed to it

¹ These conclusions are entirely in opposition to the experience of E Tree (Despendent's Quadques Drogues Actives), who found hendane to contain alkaloid in genetic quantity in the leaves, next in the first, then in the roots, and lastly in the stalk. The substance was first exhausted with pertoleum spirit to fee it from fat, then diads, finely powdered, and extracted.

F. Ransom found 0.58 per 1000 of pure alkaloid in the seeds of bennial henbane grown at Hitchin Henbane seed is used in Germany for the preparation of the alkaloid

Farr and Wright (Pharm Jons, [3], xxu 255) have proved that practically the whole of the alkaled of hembone is contained in the functions. From 100 cc of timeture (corresponding to 125 grammes of the substance), prepared from different parts of the plant, they obtained the following weights of alkaloid—

	From 100 c c Tincimo	of Plon 1	ou parts o stance
Dried leaves, average,	. 0 0103 gram:	mes. 0.0824	per cent
Recently dued fresh leav	s, 0 013 ,,	0 104	
Seeds,	. 0 015 ,,	0 120	,,,
Root-bank,	0 020 ,,	0 160	,,

From stransnum seeds, J D A Hartz (Pharm Jour [3]) xr 203) obtained 0 167 per cent of alkaloid, by extracting the fut from the dred substance by petioleum spirit, then removing the alkaloid with proof-spirit, and proceeding in the usual way Farr and Wright found from 0 16 to 0.24 per cent of alkaloid mastranonium seeds E S 6-hm id to found, m four samples of stamonium seed from different sources, 0.25 0.37, 0.05, and 0.20 per cent of total alkaloids. From 50 to 70 per cent of these consisted of pure stroptine melting at 115° G. The remainder, which was much incre difficult to cytestallies, consisted of phycosynamic, and probably other bases and their decomposition-products. But the relative proportions of the alkaloids are probably very varieble, as

with family aesistic relation of 2014 of C Tune alcolor we distilled off, and the resemble liqued listed of The filter two purposes of the contract of the contract of the contract of the contract purpose of the contract of the contract of the contract of the contract alkines with position of the contract of the contract of the contract therefore. The following figures, by Theory, represent the percentage proportions of alkalonds obtained from the chies distantial or

	Plan	destftu	to of Bio	wers	Plant in Flower Plant in Fruit							
Part of Plant	Пусво	albus	Пусво	niger	Hyosc	albus	Hvosc	nigor	Нуово	albus	Hyonc	niger
	1808	1889.	1868	1869	1908	1800	1863	1809	1868	1800	1808	1800
					_		-		_			
Seeds,	j .	١.			ă .				0 162	0 172	0.075	0 118
Leaves, .	0 588	0 469	0 154	0 192	0.859	0.359	0 117	0 200	0:211	0 158	0 005	0 110
Stalk, .	0 012		0 070	0 017	0 038	8300	0-032	0 030	0 027	0 029	0 000	0 010
Reet, .	0 128	0 176	0 027	0:080	0 148	0 262	0 127	0 128	0 100	0.088	0 028	0 056

Ladenburg found hysesyamune to piepondeaste, and Schutte found that both fresh and old stamonum seeds yielded distilly hysesyamune, with small quantities of ready-formed notopine and scopolanime. A B Lyons (Manual of Phaimaceutical Assayian) found in five specimens of stramonum seed proportions of allehold (itented by Mayer's solution) runging from 0.45 to 0.55 per cent, the extractive matter yielded to strong alcohol by the same samples varying from 3 3 to 7 5 per cent. In eight samples of stramonum leaves, Lyons found from 0.4 to 0.02 per cent of alkaloid (iterated), and from 1.9 to 2.6 3 per cent of extinctive matter yielded to spinit of 6.5 per cent. Far and Wright extracted from 0.12 to 0.22 per cent of alkaloid from stramonium leaves

R K or des ("Inaugural Dissertation," Dorpat, 1888) found the following percentages of alkaloid in the sources stated -

R. Kondes has also published the results of analyses of a large number of estracts of belladonna, henbane and stramonum. His figures show the yield of extract, the percentage of water and alkaloid in the preparation, and the proportion of total alkaloid obtained in the extract

Dunstan and Ransom (Phonm. Jou. [3], xvi. 1777 found the alkindois in mue commencial specimens of solid extract of belladoma root to vary from 165 to 445 per cent, the water ranging from 160 to 216 per cent. They state that the extract contains, beautes stropine and hyperyamme (and possible timess of another stakenois), the fluorescent substance called drystartops cade (fagge 262) and "much dextrose". This observation as of interest in relation to the assumption of Sc Av so is singer (Pharm. Zet. 1888, page 101), that a gamuine extract of belladoma leaves contains so unbistance capitale of reducing Pathing's solution at a temperature of 60°-70° C, any reddash turbulity or prespitate being probably due to sophistication with dextin or the extract of delemans or tanasamin. Analyses of vanous extracts of belladoma have or tanasamin. Analyses of vanous extracts of belladoma have been published by J. C. Um ne y (Pharm. Jon., [3), xxii. 364).

L van Itallie, recommends that the extracts of belladomna and henbane should be assayed by treating 5 gmanuse of the sample with 10 drops of dritte sulphure and (1 20), dritting with water to 50 cc, and magarating for some hours. Twenty-five cc of a 10 per

Also analyses of extracts of commum, cheledomium, acomite, nux vomice and physicstigms

eent, solution of lead acetate is then added, and after allowing the precipitate is estit the luquid is passed through a dry filter. From 50 c o of the filtrate the excess of lead is precipitated by 10 c of diuluis subplante acid (110), and the luquid agum filtered. From 50 c o of this second filtrate the akalonda are thun liberated by ammonia, extracted by three agritations with chloroform, the early evaporated, and the rendutal alkalonds dissolved in sput and trirsted with centuronal soil.

COCA ALKALOTUS!

The leaves of Esythrocylon coes and alled species contain a number of closely-alled alkalouts, all of which appear to be detrustates of cegonine, $C_{\rm BL}NO_9$ a base which Einhorn and Hesse regard as a derivative of tetrahydropyridine (pages 106, 164), and to which they assign a constitution expressed by the following formula—

$$_{\mathrm{CH_{2}}}\left\{ \begin{smallmatrix} \mathrm{CH_{2}CH[CH(OH)CH_{2}CO\ OH]} \\ \mathrm{CH^{2}CII} \end{smallmatrix} \right\} \mathrm{N.CH_{8}}$$

According to this formula, ecgonine is methyltetrahydropyridyl-\$\beta\$-bydroxypiopionic acid. When heated with baryta, it splits into carbon dioxide and isotropine, and hence may be regarded as isotropyl-carboxylic acid.

The relation between eggonine and isotropine (and therefore between cocaine and stropinc) is equally evident from a comparison of the formula of their respective anhydrides.—

The hydrogen of the hydroxyl-group of eegonine can be substituted by acetyl, benzoyl, cinnamyl, and other acid-radicals. Thus —

Benzoyl-ecgonine, Me C₅H₇N CH(O.C₇H₅O) CH₂ COOH Cinnamyl-ecgonine, Me C₅H₇N CH(O C₉H₇O) CH₅ COOH

The author is indebted to Dr B. H. Paul and M. A. J. Cownley for perusal and correction of this section.

² Upwards of eighty species of the genus Erythroxylon are found in Biaxil (Pharm Jour, [3], vm 507), but most of these species, other than E co.a, yield very little occanne (Pharm Jour, [3], xix, 70)

By heating those compounds with alkyl indules, the corresponding esters may be obtained .—

Methyl benzyl-ecgonine or occaine is the most important and characteristic of the bases of occa Methyl cunnanyl-ecgonine occurs occasionally, in small quantity, in the broad-leaved South American coos, and regularly, and sometimes in considerable quantity, in Taxillo coos

When dibsate ands teach on segmine, bothes of more complex constitution useful One of these (the methyl-nestr of a substance polyment, with ennanue said, called by Hesse cocate acid, $C_{18}H_{20}O_{2}$, and by Liebermann y-isatropic or truxillor acid, is the cocamine, $C_{18}H_{20}N_{20}$, of Hesse, and the isatropyl-occains or α -truxilline of Liebermann. The next higher homologue of cocayl-economic methyl-sets also appears to cavit in coca, as also the corresponding derivatives of its co-ocain a cid. 8

The following is a list of the bases hitherto detected in coca leaves The amorphous base to which Hesso gave the name of coolaid in his been proved to be a mixture; and the volatile base called hygrine by Lossen has not since been obtained

Isomerides of cocamine and homoeocamine probably exist in coca, as H c s s \circ has isolated from the products of hydrolysis is \circ

cocaic and homo-isococaic acids. Similarly, Liebermann has isolated two isomers of cinnamic acid, isocificant anit and allocinnamic acids, from the products of the decomposition of coca bases.

With the exception of oegonine and anhydro-eegonine, all the boties in the foregoing list are saponifable, splitting up when heated to 80°-100° with hydrochlone acid, or when boiled with alcoholic potash, according to the following equations

$$\begin{cases} a & C_{12}H_{21}NO_4 + H_2O = C_{10}H_{12}NO_4 + CH_4O \\ \text{Occurs} & \text{Demonyt-expension} \\ b & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_1H_4O_3 \\ b & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_1H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ 2 & C_{10}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_1H_4O_3 \\ \text{Consumpt-expension} & \text{Demonyt-expension} \\ b & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_2H_4O_3 \\ \text{Consumpt-expension} & \text{Demonyt-expension} \\ c & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_2H_4O_3 \\ \text{Consumpt-expension} & \text{Demonyt-expension} \\ b & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ c & C_{12}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_3 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonyt-expension} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonyt-expension} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonston} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonston} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + H_2O = C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonston} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonston} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + C_2H_4O_3 \\ \text{Demonston} & \text{Demonston} \\ c & C_{11}H_{12}NO_4 + C_2H_4O_3 \\ c & C_{12}H_{12}NO_4 + C_2H_4O_3 \\ c & C_{11}H_{12}NO_4 + C_2H_4O_3 \\$$

From these equations it is evident that the simpler bases of coca are decomposition-products of the natural alkalouis occame, occamine, homococamine, and cinnamyl-ecgonine inethyl-ester (cinnamyl-eccame), all of which readily undergo hydrolysa with formation of egg on in e, me thy all all ohol, and an arouns it a aid. Benzoyl-pseudottopius differs from the other bases of coca by redding no methyl alloud on hydrolysa.

It is evident that the mixed alkalois of cook will consist of the various natural bases in undefinite proportion, contaminated by the products of their decomposition. Hence the separation of pure occaine from the occazioning bases is very isolations of the difficulty has been overcome by I is b or in an n and G is allows of the whileston of the valueless and troublesome among which allows of the whileston of the valueless and troublesome amorphous bye-products, which are to be had in considerable quantity. The process consists in boiling the mixed bases with hydrochlore and, whereby they all suffer hydrolysis, with formation of c_0 on in c_1 and thus base forms the satirity—pout f or the subsequent synthesis

of cocaine by Einhorn's method (Ber. xxi 3335) Thus by pa-sing dry hydrochloric acid gas into a solution of ecgonine hydrochloride in methyl alcohol until the solution has become cold, and then boiling the liquid for an hour under an inverted condenser, the hydrochloride of ecgonine methyl-ester is formed, which on concentrating the alcoholic solution crystallises in prisms, melting with decomposition at 212° Cocaine is formed when this compound is heated on the water-bath with an equal weight of benzovl chloride until the mixture becomes homogeneous and the evolution of hydrochloric acid ceases The hot melted mass is poured into water, separated from the precipitated benzoic acid, and the cocaine procepitated by ammonia or an alkaline carbonate, and recrystallised from alcohol An alternative method is to convert the organine into the benzovl-derivative, and treat a solution of the latter body in methyl alcohol with hydrochloric acid gas. The artificial cocame prepared by either of these methods possesses all the characters of the natural alkaloud

Cocaine. Benzovl methyl-ecgonine Methyl benzovl-ecgonine C₁₇H₂₁NO₄, or C₅H₇(CH₈)N CH O CO C₆H₅ CH₈ CO OCH₈

Cocame is the characteristic alkaloid of coca leaves, and has recently acquired a place in the first rank of alkaloids of medicinal value It may be extracted from the plant by the usual processes. avoiding as much as possible treatment with acids and alkalies, as it undergoes hydrolysis with great facility with formation of objectionable

The s effected by Merck by treating together ecconine, benzoic anhydride and methyl iodide to 100° for ten hours in a sealed tube. The industrial reproduction of cocaine. from ecgonine has been effected and patented by Liebermann (page 272).

Cocame crystallises from a strong alcoholic solution in colourless monoclinic prisms, melting at 97°-98° C, and subliming with partial decomposition at a higher temperature

Cocame is very slightly soluble in water.1 but dissolves readily in alcohol, ether, chloroform,2 benzene, petroleum spnit, carbon

¹ The solubility of cocaine in cold water is probably near to 1 in 1800 (B. H. Paul), but is commonly greatly over estimated, owing to the case with which cocame is decomposed by hot water with formation of soluble products

2 The solubility of cocame in chloroform enabled B H, Paul to separate it from morphine, and prove a product introduced under the name of hopeine, and said to be a natural narcotic alkaloid from American hops, to be, in fact, an artificial mixture of eccame and morphine (Pharms, Jose , [3], xvi 877).

disulphide and volatile and fixed oils. It is readily removed from its solutions by adding ammonia and agitating with ether or other immissible solvent.

An aqueous solution of cocanne has a strong allkulur reaction to bitmun and methyl-orange, but does not affect phenolphibalism. The free base may be strated with the aid of either of the former undicators. An aqueous solution of cocane, if not very carried prepared and secluded from an, or preserved by an antsoptac, randly decommoses with formation of vesselable crowths:

Cocame produces on the tongue a sudden and characteristic cessation of beding, which lasts only a faw minutes. One drop a 4 per cent solution (of the hydrochloride), if placed on the tongue, soon produces a deended numbness, the effect being evanescent unless the application be repeated. Cocame also produces an interve local anasthene and blanching effect on the mucous membrane. A single drop of a 4 per cent, solution suffices to blanch the conjunctive of the eye. Anasthessa of the eye of much value in ophthalms operations, can be produced by a somewhat larget dose. Dilation of the pupil is generally produced what larget dose. Dilation of the pupil is generally produced into the system, but the mydranss produced by cocames, whether applied locally to the eye or otherwise mitcoluced into the gystem, but the mydranss produced by cocame is not invariable and is far less intense than that characteristic of atroping and its isomics.

In lago doses, cocame has marked porsonous properties. The fatal dose for dogs is from 2 to 5 grains. The hypodermic injection of $\frac{1}{10}$ grain has caused dangeous symptoms in a girl twelve years of age (see Pharm. Jon., [3], xvi. 721).

Cocame is leavo-totatory, the specific rotation in chloroform solution being about -15° .8 for the sodium ray, while the rotation of the hydrochloride in dilute alcohol is -52° 2.

REACTIONS OF COCAINE

Gocamo is precipitated from its solutions by causire alkaline, alkaline earboards and ammonia. It is almost insoluble in excess of ammonia, which is to be preferred as a precipitant. Precipitated oceanie is amophous when thrown down from strong solutions, but rapidly becomes orystalline.

¹ For various alarming symptoms produced by cocaine in dental practice, see temals by Stock man (Phasm Jour, [3], xviii 791). A resumé of the phatmeology of cocaine and its allies appeared in the Phasmaccutual Journal, [8], xxi. 161.

² If a solution of cocame salt be precapitated with caustic soda or sodium carbonate, the filtrate will be found to contain a distinct trace of benzone and resulting from decomposition of the alkaloid, but this is not the case if ammona be substituted (B. H. Paul)

Mayer's solution precipitates cocaine from extremely dilute solutions, and $A \to Ly$ on s has attempted to employ the reaction for the determination of cocaine, but with results which are wanting in exactness

Iodised todde of potassium gives a rose-coloured precipitate with a solution of 1 part of cocaine hydrochloride in 7,500 of water, in stronger solutions the precipitate appears brown, and under the microscope assumes the torm of black globules

Tannu produces a distinct cloud in neutral solutions of contine containing 1-25,000, and a distinct precipitate with twice disproportion. Piero and produces in strong solutions a yallow precipitate, najudly becoming cystalline, and appearing under unicroscopic in sheaf-like forms. Phosphouolythic and produces a fanti tribulty in solutions of 10-50,000, and a distinct cipitate with 1 12,500. Phosphotungstic and gives a golatinous with recentrals, soluble in ammonia.

Platine chlorde produces at once, m solutions of cocane hydrochlorde containing 1 400, a yellow precipitate consisting of planness needles, mostly of stellate pattern. In solutions of 1 600 most of the crystals resemble carpet-tacks, consisting of short, well-formed prissus, with a single branch from the centre, joined at an oblique angle and tapening to a point. The characters of the chloroplatinate dustinguish cocame from the amorphous bese associated with it in coca-leaves, the platinum sail of which is for less solvible in water, and crystallises in coestic-like for ins no contrasting strongly with the feathery appearance of the cocame sail:

Cocame aurochlorade is precupitated on adding auro collorade to a solution of cocame hydrochloride. In solutions containing 1 3000 an immediate precupitate is produced, which appears under the microscope in forms seembling ferm-fronds, generally with a state that arrangement. In solutions of 1 12,000 similar crystals form after a short time. "Cocambine" sunochloride forms muniter pissants crystals, having a microscopic appearance quite difficient from that of the cocame said (A B Ly on a, Amer Jour Pharm, In; IN 51).

According to Lerch and Scharges, if a drop of ferric chloride be added to a solution of cocaine and the liquid boiled, an intense red colour will be developed "owing to the formation of benzoic acid" Benzoyl-egomine also gives the reaction

Potassium bichromate does not precipitate cocame except from neutral solutions, unless they are very concentrated (1 25); but Metzer states that from a solution containing hydrochloric acid, chromic acid precipitates the chromate, Crufl, NO₄H₅CrO₄, in

ally, lustrous plates (compare page 287). If 0.05 gramme of occanne hydrochlorindo be desolved in 5 e or wister, and five drops of a 5 per cent aqueous solution of chrome send added, each drop produces a distinct presquistes, which immediately redissionlyes, but if 1 e a of stong hydrochloric and be now added, a heavy yellow presquiste of occanne chromate is produced II occanne be present, reduction of the chrome and will ensue. Regomme, apartems, attroprise caffene, pilocoupuse, oodmes and morphine do not form yellow precipitates with chrome end or plotsessum chromate. Quimme, quantities, endonme, enchonome, enchonome, enchonome, bydrochoume, apartems, brunes, brunes, strychime and versatrine form procipitates with 5 per cent chrome acid if the solutions are neutain, but, according to K Mctaze (Pharm Zeit, XXXV 697), occanne is singular in being precipitated only after addition of hydrochlore acid

F Gresel (Pharm Zett., 1886, page 132) has observed that cocaine permanganate is very stable compared with the coriesponding salts of the majority of alkaloids. Hence, if 1 centigramme of cocame hydrochloride be dissolved in one or two drops of water, and about 1 cc of a 3 per cent solution of potassium parmanganate be added, a purple-violet crystalline precipitate of cocaine permanganate is produced, the supernatant houid acquiring a purple-violet tint A B Lyons recommends that a strong solution of the cocamo salt should be used, and the permanganate employed in decinormal solution (3 162 grammes per htre) The precapitate is unstable, and decomposes in a few hours even at the ordinary temperature, leaving a brown hydrated manganese dioxide If the liquid containing the precipitate he heated to boiling decomposition occurs at once, but without the production of any peculiar odour But if examined under the microscope when first thrown down, the precipitate is found to consist, wholly or in part, according to the strength of the cocaine solution, of translucent, violet-red, rhombic (nearly rectangular) plates of great beauty, often grouped together to form roseites A 5 per cent. solution of cocaine gives a copious precipitate at once, and a 2 per cent solution after a short time; but with a 1 per cent solution the crystals only form as evaporation takes place

The behaviour with potassium permanganate serves to detect as admixture of methyl cummyl-engonine and certain other imputies in cocame hydrochloride. The presence of these causes an immediate technical of the permanganate in the cold. The first drop or two of the reagent produces a brown discoloration, while the presuptate thrown down by a further addition is more one brown, instead of a dastinet viole-pumple or red. If a limited quantity of the reagent be employed, and the hydrochloration is the contract of the present and the production of the reagent be employed, and the hydrochloration is the contract of the production of the reagent be employed, and the hydrochloration is the contract of the production of the reagent be employed, and the hydrochloration of the production of the production of the reagent be employed, and the hydrochloration of the production o

boiling, in presence of impurities a distinct odour will be developed in some cases resembling that of bitter-almond oil, and in others like that of citude cocaine (A B Lyons, Amer John Flaarn, 1886, page 240). The behaviour of other alkaloids with potassium permanagnates is described on page 144.

According to F da S11va (Compt. Rend., cx. 348, Pharm Jour., 13] xxx 163), when treated by Vitali's test for atropine (Jong 257), even a munte quantity of occusion (0 0005 gramma) developes a distinct and peculiar odorr, recalling that of properties of the produced by coherence of the produced by coesine A C Star k (Pharm, breame, and veratime develop slight odours which cannot be mistaken for that produced by coesine A C Star k (Pharm Jons, [3] xxx black has confirmed Da Silvá's statements, but considers the odour scarcely distinctive enough to ender the test completely reliable.

SALTS OF COCAINE

Cocame Hydrochlouse Hydrochloused Cocame C₁H_mNO_mCuI This sait, which is seadly prepared by neutaisang cocame by hydrochlous ead, crystallases from alcohol in short pursus malting to A B. Lyons, 9 8 per cent of wate, while those from the alcohole solution are shiptones. The sait is not hygroscopic, but is soluble in less than its own weight of water, forming a thick syrupl right if it is readily soluble in spirit, but with less facility in absolute alcohol, chloroform, and amylic alcohol, and is pracically insoluble in either, perfections spirit, and fixed and volatile oils. Ether precipitates cocame hydrochloride from its solutions in absolute alcohol ² and chloroform.

Cocame Hydrobomide, BHBr, 2H2O, crystallises readily from its aqueous solution in transparent prisms, stable in the an

Cocarne Acetate is readily soluble in water. It is difficult to obtain it in a crystalline condition, as acetic acid is given off during the evaporation of its solution

Cocaine Oleate readily crystallises, and is soluble in oleic acid and fixed oils

Cocaine gives crystalline salts with sulphuric, boric and oxalic acids. The citrate is hygroscopic, and crystallises with difficulty.

¹ Stockman (*Pharm. Jour*, [8], xvn 862) gives the solublity of pure occaine hydrochloride in chloroform, absolute alcohol, and amylic alcohol as 1 in 48, 1 in 34, and 1 in 70 respectively, but B H. Paul does not find such large proportions of solvent necessary.

Geomes Benezata, C., Ha, NO, C., H.; O., may be prepared by mixing molecular proportions of occanne and benezoe acid. It is a very soluble sait, obtainable with difficulty in accedial crystale, the solution usually drying up to a gunmy mass, which gradually acquires a crystalline structure. A sample of commercial occanne benezota of French origin was found by B II Paul to give no precipitate of occanne with ammonia, and no beaucon said with hydrochloric acid. It comessed of be no oyl-e cgon in a (Pham Jour, [3], xri, 317). According to A Big no in (Pham Jour, [3], xri, 317). According to A Big no in (Pham Jour, [3], xri, 201), the ansesthesia produced by a 5 per cent, solution of occanne bemoste lasts during four consecutive hours, and is not preceded by the sensention of pan produced by the hydrochloride

Examination of Commercial Cocaine and its Salts

The absolute purity of cocains and cocains salts intended for medicinal use is essential, as various undesirable and even dangerous symptome are produced by certain impurities hable to be present.¹

Orude Cocaine has for some time been manufactured in South America for export to European markets in place of coca leaves. which have been found hable to deterioration in transit B II. Paul (Pharm Jour, [3], xviii, 782) describes it as a white or yellowish pulverulent substance compressed into thin cakes. It contains not only earthy substances, sodium carbonate and lime salts, but also a waxy substance and traces of petroleum manufacture has probably been effected by extracting the coos leaves with petioleum spirit, washing out the alkaloid with an acid, and then precipitating it with lime or sodium carbonate. It is represented as containing from 80 to upwards of 90 per cent. of alkaloid, but the proportion of crystallisable cocaine present varies considerably, in one instance not exceeding one-half of the total alkaloid present (85 per cent). The remaining portion was precapitated on adding ammonia to its solution in hydrochloric acid in oily globules, which after a time collected at the bottom of the liquid as a viscid semi-transparent layer, which ultimately became more or less crystalline In all cases the liquid remained quite

¹ The characters and texts for comme hydrechlorabe gives in the British Phermanogene of 1885 are inalequates, and in sevent reports; growth processors in the first issue, it was meanestly described as teadily solubles on their, whereas in fact it is practically, if not also belinkly, avoidable. This maskles so corrected in the repurst, but the exposus solution is still desambled as having a butter taste, which is not a characteristic of the pure sit; and is said to yield a white prespectate with embounds of ammontum, which is not the fact. "The appears solution that the pupil of the eye B I() the argueous solution distants the pupil of the eye B I() the argueous solution distants in pupil of the eye B. (I) the argueous solution distants the pupil of the eye B. (I) the argueous solution distants the pupil of the eye B. (I) the argueous solution distants the pupil of the eye B. (I) the argueous solution of the solution and the solution are solution as the solution and the so

milky for a considerable time, in this respect presenting a marked contrast to the rapid clearing of the liquid, which takes place when pure cocaine is precipitated from the solution of its hydrochloude

The analysis of a sample of crude cocaine by E R Squibb showed.—Mosture, 3:25 per cent, residue insoluble in ethei, 5 25, impurity soluble in ether, 0.50, pure alkaloid, 89 94, and loss, 106 per cent Gou, See Chem Ind. viv. 734, 1013)

A convenient method of purifying cocaine is to recrystallise it several times from strong alcohol, and, when a certain degree of purity has been attained, precipitate the base from its solution in 10 parts of strong alcohol by addition of 5 measures of water

Paul and Cownley have pointed out that the solubility of a sample of cocaine in petroleum spirit cannot be ielied on as a proof of its purity, since cunnamyl-cocaine behaves similarly.

John Williams (You-Rook, Planm, 1887, page 502) proceed to punity and assay commercial cosme hydrochloude by desolving it in the smallest possible quantity of absolute alcohology gr 0.7893, and adding to this solution as times its measure of day either, when the cosme hydrochloude is prospitated in a fingly-drivible but peticety cystalline condition. Unfostumately, as pointed out by B II P a n l, the hydrochloude in the amoughous bease and of beargely-degomes are prescribed under the same conditions, and hence the method is useless for the assay of cude column hydrochlorider for the elimination of impurities, though serviceable for unproving the appearance of a pure salt and converting it this on convenient form for use.

Cocame hydrochloride should be perfectly colouries, and soluble in water to a perfectly colouries solution, which ought be a salesiblely neutral to litima-paper. The solution of the pure sell keeps family well, but in presence of common impurities to decomposed with great facility. In the day solid state, occame hydrochloride undergoes no change by keeping I to ought to be perfectly frei modour, but as sold it not unfraquently retains the odour of a solvent used in its preparation, or has a peculiar butyric or may small, or even a distinct benzoic odour. In any case, a sample having a distinct odour must be regardled with sunsacon.

Pure cocame hydrochloude is always distinctly uystalline, though much of the commercial active presents on amorphous or granular.

1 Paul sids that it is a mutake to attempt the particular or the product of the produ

Williams' method.

appearance The tendency to crystallise is so marked that B. H. Paul (Pharma Jour. [3], Nrui. 181) regains an amorphous condition, or even difficult crystallisability, as an indication of the nessence of impurity. Paul states that on dissolving 5 to 10 grains of a pure sample in I dinchin of water and rapidly evaporating the solution (in a glass basin) on a water-tath, the dry resultue obtained will be white and opaque, presenting a indiating crystalline structure, while in the case of an impuse mixed salt the resultee will be mixed to less yellow, thanshocats, and of a guminy or resmod character.

The most definite test for the purity of cocame hydrochloride; as and by A utrich (Bw, xx 10) to be the optical activity, it dilute alcoholoc solution, at 20° C, the specific rotatory power is $S_{z} = -(62^{\circ}18 + 0.15882)$, where $S_{z} = -(62^{\circ}18 + 0.15822)$, where $S_{z} = -(62^{\circ}18 + 0.1582)$ is a considerable of parts by weight of absolute slochol with 9 parts of vates) in 100 volumes of the solution, and c is the weight of hydrochloride in 100 volumes of the solution. When g=0, or, in other words, the solution is aqueous, $S_{z} = -62^{\circ}2$, when $g \approx 100$, $S_{z} = -68^{\circ}0$, when $g \approx 100$, $S_{z} = -68^{\circ}0$, when $g \approx 100$, $S_{z} = -68^{\circ}0$.

The characteristics of cocaine hydrochloride should be, according to Be ck urt is, that it should give a clear and colorives solution in water; leave no resulte on ignition, give a colorides solution in concenhated sulphune acid, when dissolved in the proportion of 0.20 gramme to 1 cc, that a concentrated aqueous solution should be absolutely neutral (to litume), not immediately reduce potessum permanganate and when heated with the latter reagent give off no colour of bitter-almond oil

The Geman Pharmacopova (1890) prescribes the following tests for cosanies hydrochloride —— OI gamme is dissolved in 5 cc of water, and 3 drops of diluted sulphinic acid added This solution should be coloured viole by 1 drop of a 1 per cent solution of potassium permanganate, and if kept in a closed vessel the colouration should but slightly decrease in half an hour. One of sulphinic or mine acid should dissolve 0 1 gramme of a cocaine salt without colouration.

The following test is due to H M ac lag an (Amer Drug, 1887, page 22, Pham. Jour, [3], xw 680 — One graus of cocanie hydrochloride is dissolved in 2 ounces of water, 2 drops of strong ammons are added, and the walls of the containing vessel rubbed from time to time with a glass rod, in a quarter of an hour a good crop of glastening crystals separate. When the occurse is not very pure the solution remains clean, or else deposits only a small crop. With a good sample a dense precipitate is produced either at once or on starring, and soon acquires a crystalline condition,

the liquid lapidly clearing. When the cocaine contains more than 4 per cent of amorphous alkaloid the solution becomes milky

B H. Paul (Pharm Jour., [3], xviii 783) has pointed out that the precipitate of cocains produced in Maclagan's test icdissolves if left for a long time in the ammoniacal solution, owing to its conversion into the soluble base benzoyl-ecgonine He describes a quantitative application of the ammonia test (using a 2 per cent. solution of the salt) which, in the case of good samples free from odour and colour, will fairly indicate the purity and value, but, in the case of bad samples, regard must also be paid to the character of the precipitated alkaloid. This is done by adding the ammonia gradually, with constant stirring, as long as a crystalline precipitate forms and the liquid clears promptly When the precipitate begins to form clots which adhere to the sides of the beaker, and the liquid remains milky, the precipitate already formed is separated, and the amorphous precipitate produced on further addition of ammonia collected separately 1 The following results were obtained by B H Paul by the examination of commercial cocaine hydro chloride by the above process -

		Ammonia Precipitate, per cent					
Sample Number	Water, per cent	On Sample	=On Dry Salt				
1	90	85 G 84 S	86 S 84 7				
8	50	H 84 0	84				
4	1-00 48	83 0 83 6	84 00 82 95				
6	1:19	81.35	82.88				
7 8	43 9 47	81 04 74 9	81.40 82.75				
		(Caget 664)					
9	2 00	6 Amorob 122 (80 2				
10	0.57	Cryst 43 28 Amorph 84 98	78 06				
11	2 08	(Crest 417)	75.5				
12		Amorph 317 }	1 """				
12	**	1 602	,				

The ammona promptates from the first eight of these samples were perfectly cystalline, without any trace of stackness, they deposted rapidly, and left the supernatant iquad quite clear and hight. In the case of samples 9, 10 and 11, a considerable proportion of the case of a sumples stacky nature, quite different from that obtained from a pure salt. No 12 was so impure that was innossible to effect a fractional proemtation quantitatively.

¹The amorphous alkaloid when freed from colouring matter is a clear yellowish transparent substance, resembling thick canada-balsam, and the hydrochloride forms a varinsh-like mass that cannot be reduced to powder. Paul states that the principal impurity in the last four samples was undoubtedly the hydrochloride of the amorphous alkaloid associated with occame in occs leaves (see page 287), the salts having been probably produced by evaporating the solution of the mixed bases in hydrochloria cand, and it is questionably whether the presence of this amorphous base should be tolerated in a product which unrouts to be "occame hydrochloride".

Decomposition-Products of Cocame.

librarout_scorning C₁₃H₁₀NO₂, an C₂H₂₅N(O C₃H₂O),CO OH.

This base may be prepared by the action of benzore analydrinds to benzore chlorido on segonme, and is also a product of the action of acade or water on occann. Hence it occurs as a bye-product of the manufacture of occanne. Hence it occurs as a bye-product of the manufacture of coatine. If the more than one molecule or prepared by gradually aduling a little more than one molecule of benzore analydride to a host saturated aqueous solution of one molecule of segonme, and heating the mixture on the water-bath for about a hour. After cooling, the product is shaken with ether to remove unchanged benzor analydride and acid, and the residual benzoyl-segenme washed with a little water to dissolve unaltered econome. The yield is about 80 per cent of the econome pulpoyed, and an additional quantity can be obtained by concentrating the mother-liquor and again treating it with benzoralmydriale.

Benzoyl-egonne crystallises with 4H_sO in transparent, flat, timestre priss, resembling ammonium contate, which meli at a variable temperature ranging from 87°–140° When fusion occurs at the lewer temperature (as happens when the heart rapidly applied), the substance resolutions on further hearing, and melis again at 190°, turning brown at the same time

Benzoyl-egonine is sparingly soluble in cold water, but readily in hot water, alcohol, and dilute alkalies and acids. It is almost insoluble in ether

The acetate and sulphate of benzoyl-ecgonine crystallise in pulsus BHAuCl₄ forms small, yellow, anhydrous scales, seluble in alcehol but only sparingly so in water

When heated with alkalies or with hydrochloric acid to 100° in scaled tubes, the base is decomposed into benzoic acid and ecgonine By treatment with methyl lodde it yields cocaine

¹ Benoyl-orgonine is easily modined by heating cosmic with about 20 parts of water in a closed tube. The cocaine malts at about 90°, but gradually dissolves on maintening the temperature at 100°. The change is facilitated by agitation, and in about twelve hours a clear solution is obtained, which is only family and if pure occurs was employed.

ECGONINE, 283

Benzoyl-egomine does not appear to have much, if any, assentients effect when applied to the eye, and exerts only a modurate dilating action on the pupil R Stockman states that it is very initiating to the mucous membranes, and when injected sub-cutaneously produces tetane spassma In many respects its action resembles that of cassens, but paralysis of the sensory nerves is guite absent (Pharm June, 13, 12, vs. 898).

ECCONINE C₉H₁₆NO₃, or C₈H₂₈N(OH) COOH (See also page 270). Ecgomme us obtained, together with bemzon said and methyl actionly, by the third occurrent with concentrated hydrochlorus extend 100°m scaled tubes (page 272) \(^1\) Also, when occurre or its hydrochloride us heated with 20 parts of water and 10 of baryta to 120°m scaled tubes, it is decomposed occording to the equation—

 $C_{17}H_{22}NO_4 + 2H_2O = C_7H_6O_9 + C_9H_{16}NO_9 + CH_4O$

The actual products are methyl alcohol, butum benzeate, and a compound of barum benzeate with the barum compound of econome, C2Da(ChI_kOb_2)+Ba(Db2_h-2+BLO), which forms slender pusmate needles, very soluble in water and alcohol, but only slightly soluble in teller. This compound is a convenient source of econome. On subjecting it to dry distillation it yields an is a trop in c, the chiroplatinate of which forms bulky, omage-red, deliquescent rystals containing (ChI_kOb_1)I_tPic(I_s).

Ecgonne crystallises from absolute alcohol in monoclime pusms containing I aqua, which melt at 198°, or, after drying at 140° to expel the water of crystallisation, at 200°. Ecgonne is very soluble in water, sparingly in absolute alcohol, and insoluble mether I than a slight bitter-week taste

When ecgouine is heated with moderately strong sulphuric acid, neither carbonic oxide nor formic acid is found, but a base is produced which bears the same relation to ecgonine that ether bears to alcohol. It unites both with acids and bases

C E Merck (Ber, xix 3002) states that ecgonine, when distilled with nearly dry barnum hydroxide, ynelds methylamine and not ethylamine as one of the products, thus agreeing with the behaviour of tropine when similarly treated

When ecgonine (or anhydro-ecgonine) is oxidised with potassum permanganate, or nitric acid, succinic acid is formed (Ein-

Lesbermann and Gressl obtain expension on a long scale by boiling the amoughous base obtained in the manufacture of cosms for about an hour with hydrochlore and The filtered solution is evaporated to dryness, the readine treated with a little aboult to money importures, and the reminal eigenina hydrochloride decomposed by sodium earborste, the liberated base being correstablised from alcohol.

horn, Be , xxi. 47), a fact which shows that the side-chain in the molecule of econome must be either in the α - or β -position

Econime contains a carboxyl-group, and hence behaves at once as an each and a base. It has a neutral reaction, but reacts with alkahes to form gummy compounds of funt alkahus exection, which crystallase with difficulty and use very soluble in water and alcohol. Exposuse hydrochloride, C,R_H,NO₀,HCl, forms trachine tables, difficulty soluble in a alcohol and melling at 246° C B,H₂H,Cl₀, after drying at 140°, melts at 226°. It is extremely soluble in water, and is deposited in orange-root pursus on advantage excess of alcohol to its aqueous solution. BHAuCl, is a greenish yellow, gummy substance, very soluble in water and alcohol.

With iodised potassium nodide, eegonine yields a reddish brown precipitate, rapidly changing to reddish yellow, microscopic tables or prisms. In dilute solutions the precipitate is formed only after concentration. In the animal system, cocaine is converted into ecconine, which may be detected in the urne by this test.

ANTIPRO-EXCONING C₂H₁₁NO₂, or C₃NI₁McGR CH COMD.

This base is formed by the action of phosphores exyclipted or pentachloude on eegonies, or by heating cocaine for eight hours of 160° with glacula eacts cast which has been saturated with prochloure and gas. It forms colourless crystals melting at 235°, solible in water and alcohol, but insolible in other, chiloron, bezzene and petroleum spirit. When anhydro-egonime is heated with water to 150°, m et by 1 am in e is blemsted. It committee directly with bramine to form a base containing C₂H₁₂B₂NO₂, the hydrochlourid of which melts at 184° The salts of analyse-egonime are crystallisable. BECI crystallises from absolute alcohol in white needlise melting at 240°-241°,

Bases allied to Cocaine.

DEERNO-COORING $C_pH_mNO_k$. Einhorn and Marquardt $(B_{\theta\theta}, x, x, x, i, 46, 9, 79)$ have found that by warming with aqueous potash for twenty-foun hours, eagonuse is convexted into a base which diffuse from ordnary eagonuse in being much less soluble in absolute alcohol, and having a much higher melting-point (254°), but eascealffy in hemp dexto-colotary.

From this dextro econine a synthetic dextro-cocaine may be prepared as a colourless oil, which solidifies on standing, and is readily soluble in ether, alcohol, henzene, and petroleum spirit

Dextro-cocame may be obtained in crystals, melting at 43°-45°,

¹ Hence it is best isolated by treating the solution of its hydrochloride with argentic exide (compare page 20). It may be purified by precipitation from its alcohole solution by ether

by treating its solution with a crystal of benzoyl-dextroecgonine chyl-cster

The salts of dextro-occume crystaliase well BHG1 is much more difficultly soluble than the hydrochlorde of ordinary occurs, and moles at 205° mstoad of 181° 5. BHNO₂ is especially characteristic 100 parts of water at 20° C dissolve 155 parts of the initiate, which is precupitated in crystals on adding intro said to solutions of other salts of the base. This behaviour distinguishes dextrocome from ordinary cooline. B_HH₂Pt(2)₀ gritalinies from bot water in yellowah noodles BHAuCl₂ crystallises from dulute alcohol in needles melting at 148°.

Dextro-cocaine was found to resemble ordinary cocaine in its physiological effects, except that local anæsthetic action commenced more rapidly, and disappeared in a shorter time

With chromic acid, potassium permanganate, and auric chloride, dextro-cocaine behaves like cocaine

COMMITTIALING, HOMOGOGAINM, or Benzoyl-econome ethyl-sest, CAGEMNO, as the higher homologue of cocume, which have it closely resembles. It is prepared by heating henzoyl-econome with eightyl coide and alcohol for eight hours at 100° It crystallines from alcohol in witerons prisms melting at 100° It crystallines from alcohol in witerons prisms melting at 100° It crystallines from alcohol in witerons prisms melting at 100° It crystallines from alcohol in witerons prisms melting at 100° It crystallines from alcohol in witerons are the chloroplatinate forms inlight yellow, thombic plates, seembling the cocume sail but more crystalline. Physicologically, homococane is similar in its effects to cocume, but is weaker and less toxic, and does not appear to be mydraine.

The higher homologues of cocethyline, containing propyl and isobutyl groups, have been prepared by similar means, and also by passing hydrochloric acid gas into a solution of benzoylecgonine in the corresponding alcohol.

CINSANTI-COGAINE C_BH_ANO₄, or C_bH_A(CH_A)(CH₅O)NO₅H₅O)NO₅. This base has been obtained synthetically by passing dry hydrochloric gas into a solution of cannamyl-ecquaine (prepared by heating economic with cinname analyzinde and where). It forms large colouriess crystals melting at 121°, and is almost insoluble in which the redding solution in alcohol, either, &c. When boiled with hydrochloric and it is decomposed neabily and quantitatively into emamine and, ecogome, and methyl alcohol BHG1 is prequistated as an oil which solutifies after a time on adding a large volume of this to a strong excluditated solution of the salt in alcohol B₁H₂PtCl₆ crystallines in miscoscopic needles melting at 217°. When treated with a cold solution of potassium permangul-occume and its salts immediately evolve a strong colour of benzildelybic Ottlera-lamond oil)

Cinnamyl-cocame has been noved to occur naturally in coca leaves from 'ranous sources Paul and Cownley (Pharm Jour, [3], xx 165) examined a sample of leaves containing 175 per cent of total alkaloid, neatly 05 per cent being crystallisable from petroleum spiril, but which, nevertheless, contained very little real cocaine On oxidation by permanganate the crystallisable alkaloid yielded abundance of benzaldelyid, and in other response corresponded with emnamyl-occaine (metalyl ennamyl-ocgomia)

Codamne a-Tuxullue C₂₈H₂₈N₂₀+H₂O This base scontained in notable quantity in Taxullo concluses: Hose stoud 0.6 per cent in leaves of a different kind, and states that East Indian conclusives, and especially those from Java, contain occuming in considerable amount. Inhebitument regards occuming as admitted with the base originally described by him as γ-1 satropyl cooning, and afferervade as a-truxilline, but Hoses contends that Lubermann's product was a mixture, of which cocamine was a lealing constituent.

Cocamme has a latter taste Hesse and Stockman found its physiological effect to be similar to that of contain, but somewhat weaker, and its sins-thetic action especially weak. On the other hand, G. Flakkon altides to "y-sactopyloceauco (cocamma) as a "deadly alkalord," and Lachemann describes it as a heart-poise which does not produce ansesthesia. To its presence as an impurity, the oc-assonably highly toxic effects of commercial cocame are not improbably due

Coommue is prompitated by caustic alkalies and ammonia from solutions of its salts, and affect exposure at the ordinary temperature in a desireator retains one molecule of water. It is readily soluble in alcehol, ethe, benance and chloroform, but differs from cocaine in being very sparingly soluble in petroleum spirit. Neither the free biase nor its salts have been obtained crystalized Repeated solution in hydrochiloric acid and representation by sode was the process employed by Laelermann to purify the cocumien from the occocuming (Sertuxliniae), which is also bitter, and produces numbness of the tongue very slowly by reason of its sparing solubility.

Both cocamine and its isomeride have been obtained synthetically. When hydrolysed by mineral acids they yield occomine, methyl alcohol, and cocaic and isococaic acids respectively.

Cocase Acid, $C_0H_8O_{20}$ or $C_{18}H_{10}O_{40}$ called by Liobermann γ -1satropic acid or α -truxillic acid, is produced by boiling

¹ The composition of eccamine and its allies has formed the subject of an embittered controversy between Liebermann and Hesse (*Pharm. Jour.*, [8], xxx. 1108, 1129, xxx. 61, 101)

cocamme with hydrochilous soid. The isomeric isocorais acid (6'-isatropic or \$\beta\$-trixillia caud) is the similar product from isococamme Cocan and melts at \$71\frac{2}{3}\$, is atsistess and oduniess, insoluble in wator, and nearly insoluble in ether, from which, however, it orgaliables in forms seembling between adults also (6-trixillic) and melts at \$200\frac{3}{3}\$. Both cocan and isococan consider a mean in the constant of the products of installication and isoparticles and in the constant of the products of installication.

BENZOYL-PSEUDOTROPINE. C.H., NO C.H.O. is a base isolated by Grass I from a narrow-leaved core plant cultivated in Java / Rev xxiv 2336). It somewhat resembles devtrococome but is ontically mactive, and differs from other coca-bases in not yielding methyl alcohol on hydrolysis for when heated with hydrochlone acid under a reflux condenser for some hours, it is completely decomposed into benzoic and and pseudotropine C.H. NO (see page 247) In this respect the base resembles ationine and the other tropenes Benzovi-pseudotromne is obtained as a milky precipitate which does not become crystalline on adding sodium carbonate to the solution of one of its salis The base may be extracted by ether, and on evaporating the solution is obtained as an oil which, when quite div, solidifies in radiating givetals melting at 49° C. It has a strong alkaling reaction, and is easily soluble in alcohol, other, chloroform, benzene and netroleum spirit BHCl, obtained by passing hydrochloric and gas into an ethereal solution of the base, crystallises in white needles melting at 271° The solution gives a bulky crystalline premutate with mercuric chloride B.H.PtCl, is a flesh-coloured precipitate, insoluble in hot water, alcohol and ether. BHAuCl. crystallises from water in sparingly soluble vellow needles melting at 208° The picrate forms fine yellow needles, difficultly soluble With potassium bichromate, benzoyl-pseudotropine yields a crystalline precipitate, instead of an oily one like cocaine and dextrococcune

Amorphous Bases of Coca.

In isolating cosaine there is found in the mother-liquous a warable quentity of a basic substance commonly known as "amorphous cocaine," while the names cocaine and cocaine of the cocain of the hardware scheme to the cocaine of the

¹ Liebreich finds that benzoyl-pseudotropme introduced into the eyes of rabbits occasions strong local anesthesia and a slight enlargement of the pupil, in this respect acting more like occaine than atropine. small resembling that of motions, and a bitter and atomate tasts. Stockman concludes that "amorphous occume" is in reality a solution of ordinary crystalline occume in hygrine, the hquid alkatoid said to have been found in occa leaves by Lassen. The simprhous alkatoid is extended from the occ. in greater or less amount by the processes now employed by manufacturers, and the presence is considered by Stockman to account for cettum, disagreeable effects resulting from the employment of occame commissing the impurity. Thus if the hydrochloride of the impure alkatoid be used to produce ansesthesis of the conjunctiva considerable intuction ensees

W. C Howard (Pharm Jour, [3], xvm 71) to a certain extent agrees with Stockman's view as to the nature of amorphous cocame He found that when the solution of the bases of coca in hydrochlone acid was completely precipitated with platinic chloride, and the liquid filtered after standing over-night, the mixed platinum salts obtained were amorphous or semi-crystalline, and somewhat light in colour. When the precipitate was washed with a large quantity of water at a temperature not exceeding 80° C, the cocame chloroplatinate dissolved, and the alkaloid could be obtained therefrom in a crystalline state. The fraction of the platinum salt insoluble in water when decomposed by sulphuretted hydrogen, and extracted with ammonia and ether, left on evaporating the ether a liquid base which thickened considerably on keeping, but in which no ciystals appeared even after a week It had an intensely bitter taste, formed an uncrystallisable hydrochloride, and a chloroplatmate contaming 185 per cent of platinum (against 19.3 per cent in the cocaine salt)1 and not affected by hot water, all which characters distinguish the base from the description of hygrine given by Lossen (Annal der Pharm., exxi. 374).

O Hease states that when working on the bases from the broad-leaved occa, separating the costuse as hydrochlorate "by a special process," and secretaining the absence of contume, the restand mixture was dissolved in dutue hydrochloric acid and the solution freated with ammonia in excess, this process of solution and representation being repeated until the precipitated dissolution in hydrochloric acid gave a solution which showed no finonscence on dilution with water, thus proving its freedom from hygrine. The precipitate, after being further washed with water at 80°C are cave a moleted mass which was stread on clear takes and dried at

Hesse (Pharm Jour, [3], xviii 71, 487) considers that Howard's platinum salt was hydrated, being in reshity the chloroplatinate of an amorthous bear somerie with cocain. HYGRINE 289

60°, by which means it was obtained in transparent, brittle, hygoscopic lamine which were nearly ussoluble in water and shirtle scopic lamine specific production was alkaline to bitumy, and periodeum spirit. The solution was alkaline to bitumy, with without effect on phenolphthalon (Phoem Jour., [3], xxm. 71, 437) When boiled with alcoholic barryin, or heated in a scaled tube with hydrochloric acid, the smorphous base yields be n z o i.e. a c i.d. and another product not very significant.

From a later investigation (that, xxx. 867), Hasse concludes that the amorphous bases from true core consist theigh of benzoyl compounds of an oily non-volatile base, together with some cocamine, while, on the contany, those obtained from Truzillo lawree consist essentially of cocamine, and the enmanyl compounds of the beforementoned oily base, and the cocamine is in each case accompanied by a base containing H. less than cocamine

A specumen of the amorphous base from coca examined by B H P au I (Pharm. Joss., xviii 784) is described by him as being pale yellow, and of the consistence of thick Canada balsam. It had a faint odour at once suggestive of benzon and butyne early, and a distinctly bitter taste, but produced no ansestator effect on the tongue until after the lapse of some minutes, and then very shift compared with that produced by occame.

HYERINI Under this name several bases have been described, which were either impuse or estabily dissumilar. The name was first applied by Losson to a liquid volatile base which has not suce been obtained. The hygrine of O Hesse (Phar m Jour, [3], xviii 438) is best prepared from the mother-liquid obtained in the preparation of "cocardine" from amorphous econies. This is treated with caustic sofa and cither, the ethereal solution separated and evaporated, and the results of stalled with water. The hygmine passes into the distullate, which is family accelfied by hydrochloric acid, evaporated to dryness, and the readule tacted with caustic sofa and ether. The ether leaves on evaporation a brown only residue, which, on treatment with dilute acets cand, deposits a brown smeary mass, which is filtered off, the solution again treated with caustic sofa and ether, and the ethen evaporated.

Hygrume thus obtained is a yellowish oily substance having an odour suggestive of that of quinoline. It has a slight burning taste, and a strong alkaline reaction on fitnus, but does not alter phenolphitaleum. It is but hittle soluble in water or solution of caustic sodd, but dissofters readily in alcohol, either and chloroform Hygrine volatilises with steam, and at a higher temperature may be dissilled above.

VOL III PART II

EHGI is crystallisable. Its dilute aqueous solution exhibits a marked fluorescence, not perceptible in a concentrated solution, and destroyed by sodium chlorids and other substances. An aqueous solution of largernic hydrochloride becomes milky on addition of caustic socia, owing to the separation of the free base in manute only globules, which aggregate after a time. Hesse attributes to hygreine the formula C₁½±½ in and the constitution of a trim ethylquinoline, but Lasleumann regards it as a maxture of oxygenated bases, which may be separated by fractional distillation. The most volatile boils at 193²-195², and has the formula C₂½±½ in No, but in not desirated with tropine (page 246). The less volatile portion of hygrine appears to contem C₁½±½, No, O, and cannot be distilled unchanged at the addinary pressure. Notice of these bases is affected by heating to 120° with concentrated hydrochlories and (βστ., xxii. 675).

Hosse points out that hygune probably does not pre-exist in coor leaves, but is a product of decomposition. He state when sound core leaves are mostened with ammouns, sinken with when sound core leaves are mostened with ammouns, sinken with when, and the sthert teated with dubte hydrochlora ead, the and luquid on dilution at first shown no fluorescence, but after a time exhibits this changed er distinction.

R. Stockman (Pharms Jose, [3], xvm 701) states that hygme axists no coaleaven in very munit quantity only, and some manufacturers never meet with it. He found it in occane mother choices given him by Messer Broward & Sons, and notably in the alcohole tuncture of frash cose leaves. Stockman finds hygme to distill very imperfectly with steam in presence of cosmiss. The whole of the statements respecting hygme require confirmation.

Shockman describes hygune as a brown oily liquid with a characteastes small A drop placed on the tongue causes a burning sensation. Froge were killed by the subeutaneous impection of hygune nuxed with water. There was considerable irritation at the place of injection, while the muscles all over the body, the bowds, and the serous membranes were studded with numerous minute hasmorphages.

Coca Leaves.

The coca leaves occurring in commerce are chiefly of two kinds,

¹ The treatment is stated to have decomposed the cocaine present, some benzoic sold passing over with the hygrime. It seems mobible that a difficultiy volatile or non-volatile benzoic to flygrine was formed. A better result would probably have been obtained by adding an alkali to the contents of the retor.

the one being obtained from Erythroxylor coca, which was the original trade-product, and the other, which is of more recent importation, derived from Jamaica and St Lucia Coca leaves contain, in addition to the ordinary plant-constituents and the characteristic alkalouls. co.e. at an n is a cit.

COCATANNIC ACID (C. J. H Warden, Pharm. Jour, [3], XVIII 985) has the probable composition C14H18O8 It forms a sulphur-yellow powder, which appears under the microscope in filiform crystals interlaced in masses. It melts at 189°-191° to a deep red liquid, and is only slightly soluble in cold water, cold absolute alcohol, ether and chloroform. In hot water it dissolves more readily, and rather freely in boiling absolute alcohol. A hot aqueous solution of cocatannic acid has an acid reaction. It yields no reaction with feirous salts (according to some observers, green). but with ferric gives a dark green coloration, and reduces silver nitiate slowly in the cold and immediately on heating, but not Fehling's solution It does not precipitate gelatin The alcoholic solution gives, with alcoholic lead acetate, a precipitate varying from vellow to orange-red. When heated with hydrochloric acid to 100°, cocatannic acid vislds a glucose and a phlobaphene. The products of potash-fusion do not appear to be characteristic. They are said to include butyric and traces of benzoic acid

G J H. Warden (Phaems Jour, [8], xviii, 1010, 1027) has beserved that oon leaves which as re him necessame said elso contain much alkalord, and suggests, with much probability, that the connex and allied alkalords of coos leaves exist in combination with coestamne said. Warden, in mine specimens of the dry leaves from plants grown in different parts of India, found from 038 to 1264 per cent) of and from 0385 to 1671 per cent of "crude alkalord" (average 0982 per cent). Warden did not succeed in obtaining a crystaline alkalord from Indian coca, but does not consider the non-crystalline character delatacts from its physiologonal activity (7)

A. G. Howard (Pharm. Jour. [3] xxx. 569) has published analyses of a large number of occa leaves from different sources. His results show that while Erythrosphor occu yelds about ‡ per cent of alkalaud, the proportion obtainable from most other species of Erythrosphor is extremely unsignificant, and in some cases the alkalad is wholly absent. In Brazil alone there are upwards of eighty species of Erythrosphor.

¹ The coca plant is a small shrub from 4 to 6 feet in height, growing and largely oullivated in Peru and Bolivis, and, to some extent, in Biazil and the Argentine Republic.

H. T. Pfaiffar (Chem Zeit, x. 183, 518; Jour Soc Chem. Leid, v. 161) has described the following process of manufacturing crude cosmia hydrochlarulo direct from coca laavas.—The diamtegrated leaves are digested in closed vessels at 70° C, for two hours, with a very weak solution of caustic sods and pstroleum botting between 200°–250°. The mass is filtered, pressed while still tepid, and the filtrate allowed to stand until the petroleum has completely separated from the squeous hquid. The former is then drawn off and carefully neutralsed with very weak hydrochloric acid, when a bulky, white precipitate of cosmis which a finither quantity of the salt can be recovered by evaporation

The drad product contains about 75 per cent, of real alkalond, besules traces of "hygrine," gum, and other matters: A repetition of the process proved that the whole of the alkalond was semoved by a single treatment. The soda cannot be substituted by lime, nor the hydrochloric acid by other send

Assar or Coca Laxves Pfenfice employs a smilar process for the assay of coca laves, 100 gammes of which should be digested in a flask with 400 e. of water, 50 ec of 10 per cent sola solution, and 250 e of petroleum. The intxture is kept worm for some hours and shaken occasionally, then stanned, the residue pressed, and the filtrate allowed to separate The aqueous liquid is topped off, and the only layer titusted with \$\frac{1}{2}\$ hydrochloro coll. The number of co vequired, multiplied by 0.042, gives the percentage of cocame in the sample. The fresh leaves contain from 0.3 to 0.6 per cent, but this proportion decreases considerably if the leaves have been stored for any length of time before being worked up

For the assay of coos, v. d. Marck (Joser Pharm, [5], xx. 500, Analyst, xv. 115), after a truel of varous processos, recommended that 50 grammes of the leaves should be mixed with 20 grammes of the leaves should be mixed with 20 grammes of calcased magness and mostsaned with a thill water, dired at 60°, and the mixture exhausted with ether. The other is distilled off, and the readule treated with 50 c. of 29 per cent hydrochloric and The solution is filtered, and repeatedly shaken with ether to remove colouring-matters. Ammonia is then added, and the cocaine extracted by shaking three times with 25 c.c. of either After standing for a short time over some fragments of calcum chloride, the ether is evaporated, and the residual alkaloul weighted

For the estimation of the cocains in coca leaves, A. B. Lyons (Jour. Pharm., [5], xiii. 197) recommends that the finely-

powdered leaves should be mecerated for twenty-four hours with cipht times their weight of a mixture of \$9\$ volumes of either with \$5\$ of ammonia. From an aliquet part of this liquid the alkaloid is extracted by agitation with acadulated vater, the etheir separated, and the alkaloid liberated from the aqueous liquid by means of ammonia and again extracted with eitheir, which is then evaporated to dryness and the occame weighed. The associated bases, being soluble in water and insoluble in ether, remain in the ammoniaced liquid. Lyons states that coor leaves do not contain mose than 0 \$5\$ per cent of cocaine, and somotimes the proportion is as low as 0 \$15\$ per cent. The leaves rapidly destrouted in value, so that in aix months they are practically worthless. The product from leaves contained the product of the product of the contained of the co

M Bignon (Limb) states that ooca leaves druct un damp weather, with frequent turning, and shaltered from dew and mosture, yield easily 0'8 per cent of alkaloid, and the finer sorts can give 10 per cent and upwards under exceptional urcumstances. Coea leaves druct in damp weather, or pressed into sacks before being completely drucd, undergo a guidual ferment which ends in the completed obstruction of the occasine,

OPIUM ALKALOIDS.

O pu um, the nature and characters of which are described at length on page 383, is remarkable for the large number of natogenised organic principles contained in it. At least number alkaloids have been isolated from opium, and the list is probably still incomplete. Most of these bodues have well-defined base properties, and the majority are possonous. Some of them, as morphise and naroctine, occur in opium is considerable quantity, but the greater number are present in very small proportion, and are entitled absent from some samples.

The following table exhibits the leading characters of the nitrogenised principles which have been recognised in optim. In some cases the basic character is very feebly marked, while certain of the alkaloids (eg., pseudomorphine, oxynarootine) are probably decomposition-products

¹ Pharm Jour., [3], xvi 267, xvii 506. Biguon states that the Indian never chews core leaves alone, but mixes them with sakes and lime, whereby the alkalord is liberated, and thus obtains the amesthetic properties and numbing effect upon the mucous membrane of the stomach which he desires

LIBT OF UPIUM BABES.

Alkatoid			Formula.	Dissoverer	Date.	Meltung-	Optical	Basic Character	Physiological Action.
Mombha		Г	1 1						
	٠		Southern	Serrences	1816	:	4	Strong	FOWERING DIRECTIC DOISON.
Codelne,	•		Ou Hail	Robiduet	1882	31	н	Strong.	Narcotle poison.
Thebatne, .		_	Challen NOs	Thibounday.	1836	183	н	Strong	Wielent tetame poison
Papaverne, .			ConHaMO.	Merck	1848	147	H	Very feeble	Tetameng , feebly narrotto
Meconidine, .	٠		On Hand	Hesse	1870	3	:	Strong	
Codamine, .	•		Cart aNo.	Hesse	1872	121-126		Strong	
Landanine, .	•		Capit and NO.	House	1870	106	н	Strong	Active tetanic poison.
Lendanosino,			Cull mNO4	Hesse	1871	89	ø	Strong	Tetanic polson.
Lanthopme,	•		CasH NO.	Herae	1870	200	,	Very feeble.	Tasteleas
Protopme,		_	CreH10NOs	Hesse	1871	202		Strong	Narotto.
Oryptopine,	•		CoH Bas NOs	T. and H Smith.	1881	818		Strong	Hypnotte and mydristic.
Ehmadine,	•		CHENO	Hesse	1865	222		Well-marked.	Not possonous
Narcotme,	•	_	Call SNO,	Derogne	1808	170	н	Very feeble	Feebly possonous.
Oxymarcofane, .		_	C22H25NOg	Brown (p 829)	1875	,	,	Peeblo	
Narreine,		•	ChaHgaNO9	Pelletier	1885	170	н	Peeble	Purely hypnotic.
Pseudomorphine,		•	Cad Has Noo	{ Pelletter and Thibounery }	1885	Not fusible.	×	Very feeble.	Not personous
Gnoscopine,			Cad Has NaOn	T and H Smith.	1878	88		Weak	•
Tritopine,		_	Cash by NgO,	Kauder	1890	182		Duscid base	:
Hydrocotarnne,			C12H15NO2	Hesse	181	8		Well-marked.	Similar to narcotine
		_	-						

In addition to the alkaloids in the above list, deuteropine, opionine, papaverosine, and porphyroxine (rage 330) have been described, but their existence as individuals as very doubtful

With one or two exceptions, the alkaloule of opium are strictly peculiar to Papeare zemniferum, withio, on the other hand, the poisonous alkaloid sanguinarine, which is present in all other papareneous plants, does not appear to exait in Papeare! Indeed, with the exception of protopine, which is probably identical with the interesting alkaloid in a cleyine, CallynOp. obtained by Eyk man (Pam-Book Pharm, 1882, p. 33) from

SANGUINARINE, CyflyNO4, is best prepared from the root of Sanguinaria. Canadensis (Year-Book Pharm, 1871, 810, 1875, 256, 1879, 201) The root is exhausted with water acidulated with acetic acid, the colution precipitated by ammonia, the precipitate dired and exhausted with other, and the othereal solution treated with hydrochloric acid gas, which throws down the hydroohleride of sangumarme (BHCl+H-O) as a scarlet precipitate, which may be purified by solution in hot water and repetition of the treatment with ammonia, ether, &c. The free alkaloid melts at 160°, and orystallises from hot alcohol in small white needles having an acrid, burning taste. Sanguinatine is a powerful narcotic person , the powder causes'sneezing It is insoluble in water, but soluble m ether, chloroform, amylic alcohol, benzene and petroleum spirit. The solutions exhibit a strong violet fluorescence without absorption-bands, and are optically mactive The salts of sanguinarine are change-red, and hence the free alkaloid is reddened by the fumes of hydrochloric acid. The precipitation of the bright red hydrochloride from the ethereal solution of the alkaloid, as above described, is a highly characteristic reaction. Alcoholic sulphune and beliaves similarly Aqueous solutions of sanguingine salts exhibit a violet fluorescence, and are precipitated white by ammoma and bright 1ed by potassic-iedide of moremy B2H2PtCla+H2O forms a bright orange aprecipitate, very elightly soluble in water.

gh CHELERYTRINE, which occurs m chelidonium and several other plants, phi pagarded by Schiel as identical with sangunarine, but E Schmidt agrees—this Naschold that the more probable formula is CashiyNO₄

⁷ CHELIONSTEIN, C.-H.R.O., + H.O., in the puncipal shkelood of the twelve shead to exist in the soot of the common cel and in a (Chekshowium sungue), and occors in several other plants in association with sangunatine or chelerythriae, (or both) Chickhoum forms coloulese monoclime crystals melling at time, so shills in adoilo, but modules motoclime crystals melling at the rise asit of chickhoum ser oscionieses, and have a very sent and multite tasts. The hydrochicula fram time crystals which require fully 800 parts of cold Chickhours in a sertiary loss, so come has used in such time the statistical Chickhours in a sertiary loss, so come to see the second problem of the statistic color of the color of the second problem.

Several other alkaloids besides those already named have been detected in Chekidonnum manus, among them being a- and \$-homochelidonine, in morphine both an ethyl and a methyl group are directly united to the nitiogen atom.

Pseudomorphine was formerly represented by the formula CyllmNO. Hesse found that the base contained a molecule of water which, when driven off, was recovered very rapidly. He therefore preferred the formula C,vH,vNO, but more recently has abandoned this for C17H18NO2 or preferably C34H28N2O2 the base having the constitution of an oxydimorphine 1. On the other hand, M P Cazenenve (Compt. Rend, 1891, p 805) has obtained a violet colouring matter of definite composition by acting on morphine with paranitroso-dimethylandine (page 75) This dve appears to be an indamine, analogous in constitution to Buidschedler's green, whereas, if psoudomorphine were derived from two molecules of morphine, the colouring matter would have contained two morphine residues, and had the constitution of a safranine (Part I page 252). Combination is not effected by means of the hydroxyl-group having a phenolic function, since codeine yields a sımılar dye.

Narodina, $C_{\rm B}T_{\rm B}NO_{\rm P}$ contains throe methyl-groups, (besides that connected with the integen,) the first two of which may be successively removed by heating the alkaloid with strong hydrochlore and, while by heating with funning hydrides and the third group may be removed, normarodine, $C_{\rm B}T_{\rm B}NO_{\rm P}$, being produced together with me thy 1 to 4d i.e.

When narcotine is heated with water under pressure at 150°, it is split up in the flist place with formation of opianic acid and hydrocotarnine (page 325) —

$$C_{22}H_{28}NO_7 + H_2O = C_{10}H_{10}O_5 + C_{12}N_{15}NO_9$$

The two products subsequently react more or less completely to form meconin and cotarnine, thus .—

$$\mathbf{C_{10}H_{10}O_{5}+C_{12}H_{18}NO_{8}=C_{10}H_{10}O_{4}+C_{12}H_{18}NO_{8}+H_{2}O_{10}}$$

(compare page 161)

Orienze aud, $C_{\nu_0}H_{\nu_0}O_{\nu_0}$ (compare page 203), forms delicate white crystals It is a soluted to meconin (page 336) by nascent hydrogen, and by oxidation with dilute chromic and mixture yields hemipline and $C_{10}H_{10}O_{\nu_0}$. By the action of soda-lime, opaque and yields methyl-vanilin, $C_{20}H_{10}O_{\nu_0}$ which when boiled

On heating pseudomorphine with acetyl chloride, a tetracetyl-derivetive is produced, a fact which indicates that the four hydroxyl-groups are still mtact, and that the hydrogen atoms lest in the formation from morphine must have been united with callon.

with hydrochloric acid gives vanillin, $C_8H_8O_8$ (Part I page 62; see also Dott, Phan Jour, [3], xiv. 641) 1

Colorance, Cig.H., NOp. is contained in the mother-liquor from which the meconin has crystallized. It forms a very soluble, yellow, bitter substance. It is a farrly strong bees, soluble manmonia and funsible in boling water. When gently heated with very children in the case of the contained and containing C., Hig.Op.

W Roser (Annales, othy 334, 559), from a careful consideration of the evidence, considers narcotine to contain the residues of opiame and and hydrocotarnine, and expresses it by the following graphic founds. It is closely related to paparerine, both being derivatives of a bointy 1-18 of unit of the

W. Rosei (Annalen, extvn 167) has obtained an isomer of narcome by treating narcotine methochloride in aquicous solution with causits code, when narcotine methyl-hydroxide is precipitated. On exposure to steam this changes into a base which possibly identical with narcotine, apparently in accordance with the equation $-C_{22}H_{23}NO_{p}CH_{p}OH + 3H_{p}O = C_{22}H_{23}NO_{p}2H_{q}O$; or perhaps the new base is an anhydro-narceine containing $C_{23}H_{23}NO_{p}SH_{p}O$

Narceine has been expressed by the constitutional formula:--

General Characters of Opium Bases.

Morphine, codenie, thebaine, papaverne, naccoline and narceine are the most important of the alkalods of opinii. The opinii alkaloids form a group of which all the members exert a more or less narcota and totanising action, but in very varying deposition. The morphine is almost purely narcotic and thebune almost purely tetanising in its action. I Morphine, odeline and thebaine almost have strongly-marked base characters. They are strongly alkaline to hitms, said afford stable salts. Papaverume, nursue and navezne, on the contrary, are very weak bases (compare users 300).

The free alkaloids of opium are generally but slightly soluble in water, but dissolve more readily in alcohol. In many instances the solutions of the free alkaloids are strongly alkaline to litmus. On the other hand, certain of them (e q , morphine, narceine, laudanme) exhibit a distinct phenoloid character, and form definite compounds with the alkalies The different behaviour of the opium bases to solvents affords a valuable means of distinguishing and separating them They are precuntated from concentrated solutions of their salts by caustic alkalies and alkaline carbonates, some of the precapitates dissolving in excess of the reagent. Most of the opium alkaloids (except papaverine and laudanosine) have a layo-rotatory action on polarised light, but the specific rotatory power varies so greatly with the solvent and the concentration of the solution that the fact has a very limited practical value Many of the opium alkaloids furnish characteristic colour-reactions when treated with strong ands and oxidising agents, which, with observations of then melting-points, crystalline form, and behaviour with solvents, will suffice for the recognition of most of them when m an unmixed state. Their separation is described on page 305 et seu

BEHAVIOUR OF OPTUM BASES WITH SOLVENTS.

The following table shows the recorded behaviour of the opinion bases with solvents. The figures are the number of parts of the solvent required for the solution of one part of the alkaloid. Apomorphics is not a natural constituent of opinion, but is formed by the delivilation of morphine, and introduced into the table for convenience of comparison. The figures are the number of parts of the solvent required for the solution of 1 part of alkaloid.

¹ Thebaine appears to be the most possences of the leading alkaloids of opinin. Paparenne appears to possess only very slight possences properties, if any.

² Codeme is distinctly more strongly basic than morphine, and a method of determining the former alkaloid has been based on the fact (page 923).

Benzens Petroleum	4	Insoluble. Insoluble	Soluble	Soluble.	12 Nearly m-	19 Insoluble, 56 slightly, codifi-	_	Soluble	Sparingly, cold readily, readily,	Sparingly Nearly in-	ghtly Insoluble	Nearly in-	25 Nearly 12-	Insoluble.	Insoluble. Insoluble.		Sparngly Insoluble Soluble
Chlore- Be		Nearly fu- soluble.	Very sol. Sol	Insoluble, Solu	Readily		_	Soluble Soluble Soluble	Solunte. Syn	Readil, Spe Soluble, Nes	Soluble Sil	Nearly m. Nes	_	Insoluble Ins	Nearly in-	Readfly	Solubie. Spa Beaufiy Soli
Biller		6100	Very sol- uble.	Insoluble	Rendily	140		Solubla, 647	a	Sparringly Sparringly,	Soluble, freshij	1880	50, cold,	Drsoluble.	Insoluble	Sparrngly.	Sparingly Readily
Amplio	Account	150, cold , 50, hot.	1	:	-	88		; ;	1				910		Nearly 12- poluble.		Incomble.
Alcohel		60 to 100, cold , 30,	Very soluble with green colour	Bearly in-	Readily	southing 10 4, 4, bot.		Readily Sparngly,	Beadily	Sparmgly Sparmgly	Sparrngly, hot.	Nearly in-	189, cold	Spermely,	300, cold , enally, hot.	Soluble.	Bendily.
Fater with	Аттопи	Slightly soluble.	Soluble, quickly turning	Somewhat	with	water Sparmgly Nearly maoluble.		Soluble	Inzolable.	Insoluble. Soluble	Shehtly	Nearly	Insoluble.	Soluble.	Solubla		Insoluble.
Water	Caustre Soda.	Readily soluble.	Soluble, torning black.	Solubla.	Shehtly	Insoluble Insoluble		Insoluble Soluble Readuly	Inseluble.	Soluble . Insoluble.	Soluble.	Insoluble.	Insoluble.	Soluble.	Soluble, m- auluble in strong NaHO	Soi , ppted by large ex	Insoluble.
Ŀ	Hot.	99	ı		81	Very spar- mgly		Soluble	ı		•		2000	Soluble.	8	1	Insolubla,
Water	COLLE	2300	Slightly soluble.	Insoluble	7.5	Insoluble		1:	Insoluble.	Insoluble	Newly 12- soluble	Nearly in-	2,000	Insoluble	8		Insolubie
Allentone		Morphine, .	Аропотріше,	Proudomorphine,	Codeme, .	Thebame, Papaverine,		Meconidus, Codamine, Laudanme,	Landanosine,	Lanthopine, Protopine,	Orgptopune, .	Rheadine, .	Narcotme,	Oxynancotine,	Narceme,	Tritopine, .	Gnoscopine, Hydrocofarnine,

		Corone-Ka	COLOUR-KEACTIONS OF UPIDE BASES.	OPERM IS	A853.			
į	1	Concentrated Sulphurse Acad	ulphure And					302
Упара	(sp. gr 142)	Alone,	On adding KClO ₃ or HNO ₃	Wich Sugar	Grago 146).	Frilide's Reagent (page 147).	Parto	
Morphine, .	Orange - red, turning vel- low on heat-	Cold, no colour or faint pink. See page 214. on heatage, arrable (page	See page 314	Purple, ch'uging		Fine violet, turning blue or dirty green.	Greenish blue.	COLO
Apomorphine,	Blood-red, or reddish-	No colour (or violet to brown)	,	No reaction.	1	Deep green, turning violet.	Rose - pink, changing to	/16-14G
Pseudomorphine,	Orange - red, changing to yellow	Cold, no colour, or ouve green, on heather, dugy green (or purple), and		Ohve, then dark green, chancing	1	Wolet, changing to blue	black. Blue,	ROIIO:
Codeme,	Yellow, not changing to		Blue on	Cherry-red,	Blue, on warm-	Durky green, changing to	No colour	ט מונ
Thebaine, .	Yellow	Blood red, tarning orange yellow, ohve-green on	8	violet, i	pare	Diood red, turning	No colour	, O,
Papaverne,	Yellow.	Cold, little change; on strongly heating, violet- hine afferments, violet-	acid alone No change.		Doll purple	88	No colour	r t O M
Narcoinne,	Red		Carmme red.	Fine maho- gany brown	On warming, punk, chang me to orange	blue and yellow 1 Pink, changing to green, No colour. yellow, and orange	No colour.	DADES
Namente, .	Yellow rapidly fading.	Brown, desolving to yellow solution (changing to dark red). If impure, rod or blue colour	No change	Not character	red and vio- let. Brown-yellow, becoming ins- hogany brown on heating 1	Brownish green, chang- ing to yellow and red- dish ! (yellow-brown to blue).	1	
		Great discrepancies gener in the descriptions of this generation	a demr in the	Loundinform of	this soundfor			

The solubility of opium bases, as of other substances, is much affected by the physical condition of the alkaloids, and to some extent by the manner of making the experiment.

COLOUR-REACTIONS OF OPIUM BASES

Several of the opium bases react in a more or less characteristic manner with potessium permanganate (see page 144).

Many of the opum alkaloids gre brillant, and m some cases characteristic, colour-reactions with mineral acids, with or without the aid of heat and the addition of oxidising agents. The colours obtained vary somewhat with the motile of applying the test and with the cavidiser employed. The colours obtained are modified in a maked manner by very slight times of oxidising agents in the sulphure and used; and hence this seagent should be scrupulossly free from iron and oxides of mitrogon. E. Kau de 1 recommends that the purity of the sulphure and should be tested by codenie, which should give no colour even on heating, while in presence of the faintest time of iron, such as may be taken up from long k-eping in a bottle of common glass, a violet coloration is produced.

The colour-reactions of the opum alkaloids are best observed in the manner described in detail on page 313 et seq

Many of the colour-reactions of the opium bases dedclassification, and such of these as appear of value are described under the alkalouds to which they rofer, but the table on page 502 shows many of the better/known reactions of the more important opium bases, according to the most rehable observers.

If a trace of narceme be evaporated with ditute subplurue acrd at 100° C a beautiful violet-red coloration appears as soon as the liquid is sufficiently concentrated; changing to charry-red by commend heating. After cooling, the addition of a trace of intra eard or a mixte produces bluesh violet steaks in the red liquid. The test which is due to Plugge (Jown Chem Soc., in 870), is said to be very delecta and characteristic With traces of morphine, codone, or papaverne the liquid remains quite coloraless, with larger quantities of either of the two former bases a faint reserved in its obtained, with thebane a greenish yallow to brown colour, and with narcotine a red to reddish brown

According to Serena (Analyst, x 149), the following colourreactions are produced on treating certain of the opini alkaloids successively with a few drops of concentrated sulphure and and a very small quantity of a dilute solution of ferric chloride, with the aid of slight heat

Alkalond	With Sulphuric And	On adding Ferris Chloride
Apomorphine,	Not changed	Violet streaks at point of contact, the blush green mass becoming light violet on heating
Codeine, .	Light violet red, despensed by heat (compare p 822)	Sky-blue
Papayerine, .	Purplish red	Colourless, on heating,
Opienine,	No coloration	Green, rapidly becoming
Narceine,	Coffee brown	Bluish green
Codnmme, .		Green-blue , at 100°, violet

The following table shows the colour-reactions observed by Hesso. (Jose. Chem. Soc., xxiv 1064) when certain of the opium bases are treated with pure concentrated sulphuric and, and with acid containing traces of oxide of iron or oxides of nitrogen. The reactions with fearire chloride are also shown.

A likedoid	With purs Su	book orund	With Aoud Oxide	containing of Iron	With Ferris	
	At 20° O.	At 150° C	At 20° C	At 150° 0	Chiorige	
Codeine,	Colourless.	Dirty green 1	Blue	Dirty groon	No reaction	
Codamine,	Colourloss	Dirty red	Intense green-	Deep violet	Dark green	
Lanthopine,	Colourless.	Brownish yel-		1	No reactions	
Landanine,	Very faint rose red		Intense rose colour	Gross, chacg- ing to deep violet	Emerald green 2	
Laudanosine,	Faint rese-red	Deep red- violet	Brownish - red (resembling schalt ni trate solu tion)	Green, chang ing to deep violet	No reaction	
Protopino,	Yollow, chang ing to red and binish red	Disty green ish brown	Deep violet	Dirty green- ish brown	No reaction.	
Cryptopine,	Yellow, chaog	Dirty green	Deep violet	Dirty green	No reaction	
Hydredotai nine,	ing to violet 2 Yellow.	Crimson - red, changing to dirty red- violet		Dirty red violet.		

² According to E. K. and der (Tharra. Joser, 183, xrun 260), if the sulphunca and be quite pure no coloration is pudied with obelene over an inesting, tota shine colorum produced if snose of iron he present: Cryptome dissolver with voice colour, changing to deep blas, and inding to greensh on standing or heating to 150°. In presence of orde of iron, cryptopine us and to describe in sulphune and with deep violetnees colour, changing to visics and deep blas, and becoming grounds on heating to 150°. The hydrochloride gives a violet colorum when this treated with ead

² According to Merok, laudanmo gives a violet colour with ferric chloride,

Hesse employs the colour-reactions of the opium bases with pine sulphuric acid as a means of grouping them, thus —

*	0 1 0 ,
Odoration at 150°.	Allalvids
Dirty dark green,	Codoine, morphine, pseudomorphine
Disty red-violet	Codamine, laudamine, laudamosine, narcotine, hydrogramine
Dirty green to green-brown	Thebame, cryptopine, protopine
Dark violet or blue	Papaverine 1
Black brown to dark brown	Narceine, lanthopine

With acid containing non, codamine, laudame and laudanosme are stated to give a dark violet colour, while narcotine and hydrocotamine react in the same way as with pure acid

It will be seen that several of the reactions described by Hesse differ in a marked manner from those recorded by other observers. As in the case of other colour-observations, the only safe way is to compare the substance under examination side by side with products of known unrity

La fon 's reagent, prepared by dissolving I gramme of ammonium selemite m 20 c of stong explinime acid, is stated by da S 11 vs (Compt. Rand, cxn. 1266) to give the following colour-reactions with the opium bases.—Codeme, magnificant green coloration, maphine, greenish blue, changing to chestnut brown, nearcions, blue, turning violet and then reddsh, with slight reddish prescriptate after long standing; increase, yallowsh green, changed to brown and red, with red precipitate on standing, pageness, blue, passing to dull green, violet and red, with a slight blush precipitate, on standing, resentiate on standing.

DETERMINATION AND SEPARATION OF OPIUM BASES.

Morphine, codeme, and thebame may be thated with case and accuracy by a standard numeral acid, using litmus or methylorange as an inductor (page 130) On the contrary, they have little or no action on phenolphthalen, the reaction with which, however, is not shanp in the case of morphine (page 311)

Papaverine, narcotine and narceine, on the contrary, do not affect litinus, and their salts may be titrated with litinus and stan-

¹ Hoses states that, when absolutely pure, paparenne dissolves in small quantities of enliptions and without coloration, its, generally, on warming a crystal of paparenne with concentrated sulphurne and, a dark blue color in produced. Dett also obtains no coloration in the cold, and the blue color on strongly insting only. A sed coloration before heating is generally due to theleanse.

dard alkal, just as if the acid were uncombaned (Plugge, Planm Jons, [3], xx 401); and the first two of them being alkaloids also evines their feebbe base characters by the fact that they are extracted by chlorroferm from and solutions. Thair salts, especially with certain organic acids (e.g., sosts, benicle), are very unstable, many of them being decomposed slowly by cold and inpully by hot water. Hence, when a compound of the alkaloid with a unuseal soid as treated with a neutral solution of acotate of socioum, or even with a slightly soul solution, the fice alkaloid sprengisted in 40,000 of narcotine, 1 in 50,000 of paparetine, and 1 in 600 of narcome, none of the other cours based on the solution of the other course.

On the foregoing and similar facts, P. C. Plugge (Analyst, zir 197) has based the following process of separating the leading alkaloids of outum. The aqueous solution of the hydrochlorides is mixed with a concentrated solution of sodium acetate, and filtered after twenty-four hours The precapitate, consisting of pure narcotine and papaverine, is washed with a little water, and dissolved in a minimum of dilute hydrochloric acid. The liquid is diluted till it contains not more than also of narcotine, when potassium ferricyanide is added. This precapitates papaverine very perfectly. After standing twenty-four hours the hauld is filtered, and the precipitate of papaverine hydrofeirieyanide either weighed as such, or washed with a little water, decomposed by dilute caustic sods, and the liberated alkaloid dissolved in dilute acid and reprecipitated with ammonia. In the filtrate from the precipitate produced by the ferriovanide the narcotine is precipitated by ammonia. The filtrate from the precipitate produced by sodium acetate is concentrated to a small volume at 100°, cooled thoroughly, and filtered after twenty-four hours The deposited narceine is filtered off, and washed with a little water. The filtrate is mixed with a strong solution of sodium salicylate, and the crystalline precipitate of thebaine salicylate separated after twenty-four hours, and washed with a little water. dried at 100°, and weighed. On subsequent treatment on the filter with dilute sods or ammonia, till the washings are free from salicylic acid (as indicated by evaporating to dryness, and the nonproduction of a violet coloration on moistening the residue with

¹ This observation a done to P. O. Plugge (Ario, Phorra, [3], XXIV 994; Assales, XXI 197]. The reaction not only dampsame papersant, narrotine and narceine from non-plum, codeme, and thebane, but also from callina, occurse, comme, atropuse, piscoorapea, stripchame, button, quinte, cutchodine and emchonation. The cunchons bases are precapitated if the sodium section is at all alkalma.

ferric chloude), pure thebane is left. The filtrate from the thebane salicylate is acidulated with hydrochloric acid, the precipitated salicylic acid filtered off, and the filtrate repeatedly shaken with chloroform This dissolves the remaining salicylic acid, and traces of narceine and thebaine, which may be recovered by evaporating the chloroform The acid liquid separated therefrom is concentrated somewhat, made exactly neutral to litmus, and mixed with notessum throcyanate (sulphocyanule), which throws down the codeme as an acid thiocyanate. Twenty-four hours should be allowed for its complete separation. The filtrate should be treated with a slight excess of ammonia, and time allowed for the separated morphine to become crystalline. The liquid is then shaken with chloroform or ether to remove the remainder of the codeme and traces of other bases. After separation it is acidulated to dissolve the morphine, heated to 60°C, and the morphine shaken out with hot amylic alcohol, after addition of a slight excess of ammonia or carbonate of sodium. Plugge's results, obtained in test experiments, except in the separation of codeme and morphine, were very satisfactory, considering the difficult nature of the problem to be solved. But the methods are not to be regarded as having the same quantitative accuracy as those for the separation of the metals

Another method of separating the principal alkaloids of opium consists in treating the solution with an alkaline carbonate or ammonia, and agitating with benzene, when morphine and narceine are left insoluble, the remainder passing into the beazene. Much the same separation occurs with chloroform, except that pseudomorphine is left with the insoluble alkaloids.

D. B. Dott has communicated to the author the following method of separating the chief bases of opium -Treat the solution of their mixed hydrochlorides with a 10 per cent solution of caustic soda, and wash the precipitate, which will consist of narcotine, papaverine and thebaine, the alkaline solution containing morphine, coderne and narceine. On acitating the filtrate with chloroform. the codesne will be extracted; and on separating the alkaline liquid, acidulating it, and rendering it faintly alkalino with ammonia, the morphine will be precipitated, the narceine, from its greater solubility, remaining dissolved. It can be recovered by

¹ The separation of codeine and morphine by this process is very imporfect, If the solution be too strong, morphine is precipitated with the codeine, and if this condition be avoided the precipitation of the codeine is incomplete. In test-experiments Plugge only recovered 70 per cent of the orderne used. Hence it is better to omit the precipitation with thiogyanato altogether, procipitate the morphine with ammonia, and extract the codeine from the filtrate by ether or chloreform, after adding caustic sods (compare page 323).

evaporating the liquid to digness and treating the residue with strong alcohol. From the bases precentated by caustic sods, the thebases can be separated fauly well by erystallisation as acid tertrate.

Narcotme and papaverine may also be separated from thebaine (and codeme) by dissolving the free bases in dilute alcohol, rendering the bound faintly and with acetic and, and adding three volumes of boiling water, when the narcotine and papaverine are precipitated; or sodom acetate may be used as already described. Narcotine and papaverine may likewise be separated by solution in boiling water containing one-third part of exalic acid, when an acid papaverine exalate crystallises out on cooling The process should be repeated several times, and the narcotine finally precipitated by ammonia and crystallised from boiling alcohol

The following is an epitome of Hesse's method of separating the rarer oppum bases from the mother-liquous left from the preparation of morphine by the Robertson-Gregory process 1 The aqueous extract of opium is first precipitated by calcium chloride, the filtrate from the calcium meconate concentrated, and the hydrochlorides of morphine, pseudomorphine and codeme separated by erystallisation. The mother-houer is diluted with an equal bulk of boiling water, excess of animonia added, the precipitate removed by filtration and dissolved in acetic acid. The filtrate is agreed with other, the othereal layer shaken with excess of acetic acid, and the acetic solution mixed with that of the ammonia precipitate The acetic and solution is then treated with excess of caustic soda, which precipitates papaverine, pareotine, thebaine, some cryptopine, protopine, laudanosine and hydrocotainine, while lanthopine, laudanine, codamine, meconidine, and a portion of the cryptopine remain in solution. The alkaline liquid is ncutralised, ammonia added, the bases again extracted by other, and shaken out with acctac acid. The acetac acid is neutralised with ammonia, when a little lanthonine sensiates out in twentyfour hours, and the filtrate is treated with more ammonia precepitate formed is dissolved in a very small quantity of boiling dilute alcohol, which on cooling deposits white crystals of mixed laudanine and cryptopine On evaporating the alcoholic solution.2 and treatment of the residue with ether, a solution is obtained from which codamins may be isolated, either by addition of fused

¹ For E. Kauder's modification of Hosse's method, see Arch Pharm. ocxxviu 419 , and Jour, Ohem. Soc , Ix 227.

² Hesse could obtain no meconiduo from this solution, and hence concludes that it had been decomposed by the preceding operations, as he had moviously obtained it from a similar source by another process (Ann. Cham. Pharm, chn. 47; Watts' Dut Chem. vs. 883)

ealeum ehloride (which causes water, colouring-matter, and crystals of codamine to separate), or by conversion into the acctate, and this into the hydriodide

The mixture of bases insoluble in caustic soda is digested with diluto alcohol, and seetic acid added tall the liquid is faintly seid to litmus On adding three measures of boiling water, a crystalline precipitate of papaverine and narcotine is thrown down filtrate, freed from alcohol by evaporation, on adding strong hydrochloric acid, will give a meannitate of civitonine hydrochloride, but in order to avoid the conversion of thebaine into its non-crystalline isomer thebascine, it is preferable to add tartanc acid, which throws down erystalline thebasne acid tartiate. The mother-liquor of this is neutralised with ammonia, and mixed with 3 per cent of its weight of sodium bicarbonate made into a paste with water. After standing about a week, a black, pitchy mass separates, the filtrate from which gives with ammonia a precipitate which is treated with boiling benzene, the filtrato being also extracted by agritation with benzene On shaking the united benzene solution with a saturated aqueous solution of sodium biearbonate, laudanosine crystallises out, and the benzene filtered from this yields hydrocota nine hydrochloride on passing hydrochloric acid gas The portion of the ammonia precipitate left undissolved by benzene contains cryptopine and protopine. These bases are converted in hydrochlorides, and the solution treated with strong hydrochloric acid, when the protopine hydrochloride forms a horny doposit which adheros to the sides of the glass, and is easily freed from the gelatinous cruntonine salt by washing with a little water

Navvene is mentioned as existing in the liquors, but the stage at which it is separated is not stated

Morphine. Morphia

C17H19NOg; C17H17NO(OH)g, or C17H17(OH)NO OH.

Morphine is the most important of the bases contained in opisim, in which it exists in combination with sulphuric and meconic acids. The mode of preparing morphine may be gathered from the mothods of assaying onum (see also last pace)

Morphine erystalizes in transparent, colouless, trimetre prisms, which are usually very short. They contain one molecule of welct, which is given off-slowly at a temperature of 90° and more inputly; which is given off-slowly at a temperature of 90° and more inputly; and 10° C (Pharm Jons. [8], xviii 701, 801, 1xx 6, 1148, 180, At or above 200° morphine partially volatilizes, melts, and turns hown, becoming carbonised at a somewhat higher temperature

 $^{^1}$ D B Dott found the proportion of water lost to correspond more usually to $8C_{17}H_{19}NO_9+9H_qO$.

Morphine is inodorous, has a persistent bitter taste, and is a powerful narcotic poison

Morphine as nearly usefulble m cold water, requiring, according to Chastan 193, 33,353 parts at 3° and 455 at 29.1° At 42°, the solubility w 1 m 2380, and in boiling water about 1 in 460 (Year-Book Pharm, 1882, p 30). The solution has an alkaline reaction. Morphine dissolves in 30 parts of boiling or 30 of cold absolute alcohol, and in a somewhat smaller quantity of rectified algorith. In ether and chloroform it is almost mostable when in a crystallised state, but dissolves sparnigly when facelily-prespitated and amorphous. A useful solvent for morphin is a mixture of equal volumes of ether and accts either (ethyl acctate), but even in this its solutibity is limited, especially in the crystalline state. Amythe alcohol dissolves morphine sparnigly (1.150) in the cold, but when heated it as fairly good solvent for it (1.50). The alkaloul dissolves best when liberated from one of its salts in precession of anniva alcohol.

In beazene and petroleum spirit, morphine is practically insoluble, as also in volitile oils

According to Florio (Gaz Chim. Italiano, xiii. 496) 100 parts of the following solvents dissolve of morphine —

			Morphine	dissolved by 100	of Solvent
Solvent			At 167-11° C.	A1 50° O	At 78° C
Alcohol, absolute,			1 132		8 623
, 60 per cent,			0 377		2 991
m 75 put cout,			0 228		1,985
Wood spirst, .			_ 1875	8 469	
			0.203		2:247
Benzone,			0 020	1 235	***
Chiocoform,			0.040		
Ether, absolute, .		- 1	0.028		***

A. B Presecti (Jour Chem Soc, xxix 405) has pointed out the great influence the physical condition of morphine has upon its relation to solvents, and has determined the proportion of difficient solvents requisite for the solution of morphine in the crystalines, unophous, and "mascent" conditions, by the last term meaning that in which the alkaloid exists when liberated by ammonia or an alkaline carbonate from the agreeous solution of one of its saits. The following are Presout's figures:—

Dott gives the solubility of morphine in water at 15° C. as 1 in 2500

		Paris of Solv	ent required.	
Condition of the Morphme.	Ether	Chieroform.	Amylio Alcohol	Benzene,
Crystallised,	6148	4379	91	8930
Amorphous powder,	2112	1977		
"Nescent" state, .	1002	861	91	1997

Other figures for the solubility of morphine are given on page 301 Solutions of counts potash and sods dissolve morphic readily, as also do buryta and inne water, and, to a limited extent, ammonia also. Solutions of caustic alkales dissolve quantities of morphine equivalent to the bases contained in them, with the formation of instable morphinates which are decomposed by actionic acid and assume a dark brown colour on exposure to an Crystaline morphinates of potassium, barnum, and calcum have been obtained From these facts, and the bine reaction with ferric olioride, Chastaing (Jon Pharm, [5], iv 19) inferred that morphine possessed a phenoide character, and this view has been fully borne out by the later researches of Grimaux and Hesse (near 289)

Solutions of morphine are law-rotatory In alcoholic or dulute and solution, S_a is said to be $-89^\circ 8$ and $S_a - 70^\circ$ For the hydrocholoride, the value is $S_a = -100^\circ$ of $-1^\circ 14$ C. In alkaline solution, the value of S_a for morphine is stated to be $-45^\circ 2$.

Morphine is very sensitive to the action of oxidising agents, a fact which is often used for its detection (page 314 of seq.). It reduces salts of gold and silver, permanganates, ferrioranides, iodic and periodic acids, &c. The reactions of morphine with strong sulphune and nitric acids are described on proges 318, 314.

When morphine is heated with strong hydrochloric acid or zinc chloride it loses the elements of water and is converted into a po morphine, Cir. High NO. (page 319)

SALTS OF MORPHINE

Morphine dissolves readily in dilute acids, forming salis which are perfectly neutral in reaction to litmus and methyl-orange, and hence it may be intraised with accuracy by the aid of standard hydrochloric acid and either of these indicators. With phenophitalian morphine does not give a sharp reaction, but the poor for neutrality is approximately the same as if the acid of the morphine salt were in a free state.

The salts of morphine are mostly crystallisable, and are all bitter and very poisonous. They are generally soluble in water and in alcohol, but are insoluble or only slightly soluble in amylic elcohol, ether, chlorofoim, benzene, or petroleum spirit. Morphine is not removed from its acid or neutral solutions by agitation with any of the above solvents, except imperfectly by amylic alcohol.

The following table shows the formulæ of the more important salts of morphine, the percentage of morphine hydrate, the relative dose, and D B Dott's figures for their solubility in cold water (Pharm Jour., [3], xin 404, xvi 653) —

Могрдина	Salt		Formula	Morphins Hydrate, per cont	Reintyes Doss	Solubility in Water at 15° 5 C
Hydrochloris	de,	,	BHCI + SH ₃ O	80.69	1 00	1 part to 24
Sulphate,			$B_{2}H_{2}SO_{4} + 5B_{2}O$	79 94	1.00	,, 23
Acetute,			B,0gH40g + 8HgO	75 98	104,	,, 2½
Lactate,			B,C3H4O3	80-80	1 00	,, 8
Tartrate,			$B_{2s}C_4H_6\Omega_6 + 8H_2\Omega$	78 29	102	,, O#
Meconate,			B207H407 + 5H20	70 46	114	,, 84

Morphens Hydrochloruts, on Morphen Rydrochlorate, BMC1+3H_O, crystallases in coloniloss allky fibres, soluble in half its weight of boiling wates and in 40 pasts of cold neutrided spirit. It becomes anhydrous at 100° C. The commercial salt often has a buff or brownish that from admixture of resultons matters, which are detected by the brown or black colour assumed by the salt when hetsel to 130° C.

Mophuse Hydrichide, BHII+3HI,O, is obtained as a compact mass of hut-the noddles on muxing a concentrated alcohole solution of potassium noddle with a concentrated solution of morphims hydrichlorida. The product only slowly techsolves on adding more spirit, and is very spaningly soluble in water, especially in presence of potassium iodule. The hydrolnosiuds can be obtained similarly.

Morphine Sulphate, B₂H₂SO₄+5H₂O, closely resembles the hydrochlorido. It loses 3H₂O at 100°, and the remaining two atoms at 110°. It exists naturally in opium.

Morphine Acetate (see above) is a white, or faintly yellowish white, obscurely crystalline powder. It is readily soluble and crystallisable. It is partially decomposed by bothing or evaporating its aqueous solution, crystals of morphine being deposited.

Morphine Tentrate, $B_2C_4H_0O_6 + 3H_2O$, is readily soluble, but the acid tentrate, $BC_4H_0O_6$, only sparingly so. Their solutions are not precipitated by caustic alkalies, alkaline carbonates, or chloride

of calculum The tailante is best detected by precipitating the concentrated solution with potassium acetate and acetic acid in presence of alcohol (Vol I. page 467). After boiling off the alcohol, the morphia can be precipitated from the filtrate by an alkaline carbonate or ammont.

Mo plane Moconate (see above) is interesting as being the form in which inorphia largely exists in optim. When morphia hangely exists in optim. When morphia and meconic and are dissolved in absolute alcohol, and the solution is evaporated, an amorphious, hygoscopic, vary solubble residue is evaporated, in concentrated solution deposits citystals of neutral morphine meconate containing 5 aqua, even in piesence of sufficient meconic and to form the and salt

DETECTION AND DETERMINATION OF MORPHINE.

Free morphine, when pure or in the form of one of its ordinary salts, is readily detected. Its determination is easy when unmixed with interfering substances, but as it exists in opium is attended with considerable difficulties. Most of the colour-reactions of morphia are best observed by operating on the solid substance, but for certain qualitative tests and for all quantitative methods the alkeloid must bur asolition.

- A. Reactions of Solid Monphine For observing these reactions a munter fragment or crystal of the solid abloulor ents sails should be employed, and the experiment should be conducted in a small procedule has not or enceible. The readule obtained by the experiment of the solition of morphine in alcohol or amylic alcohol is well-smited for the operation.
- 1 Solid morphine bested with a drop of a perfectly nested colution of ferno chloude or iron-alum gives a very characteristic deep greenish blue colour, changed to green by excess of the regent. The colouring matter is not taken up by chloroform. The colour is destroyed by free acid, by heat, or by contact with alcohol. Pseudomosphine also gives a blue colour with ferno chloride, and codamine a dask green.
- 2 Nitric acid (142 sp gr) added to solid morphia turns it an orange-red colour, which is changed to yellow on heating, and dostroyed on adding sodium throstliphate (hyposulphita). The
- ¹ The colosaton is produced in stong solutions of morphisis, but bosones improperpile with mediante ablition J L A in this page (Phorm Jony, 13), xxin, (24) has pentied out that even in solutions far too didn't to give the scarcion, the inception may be detected by adding poissassin feirogrands, which produces a bins or groun colosaton. Armitings attributes tins-section to this sedicution of the non to the fearous state, and the reaction of this with the fearous made to form Turballial blue, but it is more probable that the fearonamic is reduced to ferroryanade, and then reacts with the fear on the comparison of the comparison of

coloration is said to be due to the formation of a body of the formula $C_{10}H_0NO_{g_0}$ which yields pierre acid when heated with water to 100°.

- 3 Solid morphine, when pure, is commonly said to yield no coloration in the cold on adding pure concentrated sulphure said, but according to Dott (Pharm Joss., [3], xn 615) a distinct, thought fanit, pink colour is produced. On heating to 150°, a divity green for 108-8-ed) colour is developed, and on rasing the temperature still further the solution becomes almost black On allowing it to cool and dulumy with wates, a greenal bile colour is proluced, which on addition of ammonia in excess becomes green.
- 4 On adding oxidising agents to the solution of solid morphine in cold concentrated sulphnic acid, the following reactions are produced 1 a After adding a drop or two of water to heat the mixture, the subsequent addition of nitric acid will produce a 10%-red coloration, changing to brown The reaction is very delicate b Potassium chlorate gives reactions similar to those with intric acid If the alkaloid be first heated with concentrated sulphuric to 100° for half an hour, and a crystal of potassum chlorate or natrate added to the previously cooled violet-red solution, a beautuful violet-blue colour is produced, which passes into a dark blood-1ed, changing to yellow c If the sulphuric and solution be heated on the water-bath to 100°, and a minute fragment of pure potassium perchlorate2 be added, a deep brown or reddish brown coloration is produced, which rapidly spreads through the liquid The colour is destroyed on dilution. L Sie bold, to whom the test is due, did not observe a similar reaction with any other alkaloid & Potassium bichromate is ieduced with production of green colour (No colour-reaction is produced if for the bichromate be substituted the dioxide of lead or manganese. Distinction from strychnine) & On adding sodium or potassium arseniate, and warming gently, a slateblue colour is produced, which on mising the temperature passes into green, then into deep blue, and finally, when the and begins to volatilise, again into dark olive-green. On diluting moderately with water, a reddish brown coloration is produced, changing to dirty bluish and green on further dilution; and on agitating with chloroform the latter liquid is coloured violet-blue (Donath). If

¹ The reactions in question have been verified in the author's laboratory by W. H. Ballaclough, and the description given in the text is in accordance with his results.

² The perchlorate must be free from ohlorate, which is ensured by heating it with hydrochloric acid as long as chlorine is evolved. The salt is then washed with cold water and dued.

sodium phosphate be substituted for the arsemate. and heat applied till acd funes appear, the mixture becomes voide, heat applied till acd funes appear, the mixture becomes voide, heat applied till acd the substitution of the substitution gives a fine voide to the substitution of the s

5 If solid morniume be mixed with from 2 to 8 parts of powdered cane-sugar, or solutions of the two bodies be mixed and evaporated to dryness, addition of a drop of concentrated sulphuric acid will produce a beautiful purple colour, changing gradually to blood-red and brownish red, becoming ohve-brown on dilution with The colouring matter is not soluble in chloroform. The test may be applied to a solution of morphine by saturating the liquid with sugar, and pouring it carefully on to some concentrated sulphuric acid, when a purple or rose-red coloration will be observed at the junction of the two fluids. Codeme gives a very similar reaction (Schneider) According to H. Weppen the delicacy of this test is much increased by adding a drop of brominewater after the sulphuric acid, this modification rendering the reaction equal if not superior to reactions 3 and 4 c. and less dependent on the purity of the morphia.

M Robin mixes the alkalord with twnes its weight of powdered sugar, and adds one or two drops of pure sulphune send, and states that morphine hydrochloride gives a beautiful rose colour, changing first to the tint of a solution of potassium permanganite, and then to volet and dark green, while codeine gives a cherry-red colour changing to violet, and narcotine a beautiful and very persistent malocant-brown colour ²

B Reactions of Morphine in solution The following reactions

¹ For convenience, this test is described here, but it seems improbable that the reaction is due to oxidation

Attorpus gives with sugar and subplume and a veolet coloration, changing to blown, restrine, a deep green, sanisam, a red colour, changing to collicate. Salicia gives a vival red. Puro accentine gives no reaction, but mixed accente alkalend as extracted from the root give a fine cherry-sid colonation, changing to custom. No esection is given by strychance, brucene, cocano, pilocarpine, cuffens, buberne, apometphine, cupremo, on the curchons bases (I F B ui n ttl.).

are yielded by an aqueous solution of the hydrochlorido or acetate of morphine —

1 On adding to a tolorably concentrated solution of a salt of norphine a Kawa caustic alkala, an alkaline extended, and or him extended, and the salt of the control of

A fauly accurate determination of morphine may be made in the absence of interfering substances, by peocriptating the tolerabily concentrated, cold, aqueous solution with sodium herabonate, allowing time for the precipitate to become crystalline, filtering, washing moderately with very cold water (prefembly saturated with mophines), drying at 100° or 120°, and waighing the anhydrous morphine, O., Ha, NO_o, when the wealth becomes constant

Instead of drying and weighing the alkaloid, the weaked preplate may be placed, together with the filter, in a moderate concesor of standard send, and the excess employed ascertained by titrain with litims or methyl-orange (not phenolphithaleus). I ac of decimomal acid neutralises 00285 geausme of anhydrous morrobine.

2 If morphia be hierated from the solution of a sail by one of the reagents mentioned above, and the hugh and surponded prequisits be at once shaken with hot empire alcohol, cold sectic either, or a mixture of equal measures of ether and acetic ether, the morphia passes into solution, though with some difficulty, and may be obtained in a free state by separating the othereal liquid, and evaporating it to dryness at a gentile heat If the hierated morphia be allowed to crystallise before subjecting it to agistation with the solvent, its solution becomes very difficult to effect.

For quantitative purposes, but anyluc alcohol should be employed, as the colvent II should be added before the alkahol as liberada, which should be done by ammonia, magnessa or sodium brearbonnies, and the agitation should be conducted immediately, and the separation and re-agitation offected without delay On evaporation of the surphe about at 100° the amylydrous morphism will remain as

¹ The sestic ether must be free from said. This may be ensured by agritating it with some sedium breatbonate before use.

a readine, which can be weighted, or the amylic alcohol containing the alkeloid in solution may be taisated by dulties standard seed and methyl-crange, as described on page 131. If desired, the alkeloid may be recovered from its amyle alcohol solution by repeated agitation with dhitte hydrochinus early, and then repering the contract of the contrac

- To effect complete extaction of the morphine liberated by magnessa, amonta, or an alkaline hearboants, awernal egistations with amylic alcohol are necessary. If ammonia be employed, sufficient passes into the amylic alcohol to vituate the subsequent determination of the morphine by tirution; while if the amylic alcohol be freed from ammonia by agitation with water, or even with built, a protinc of the morphine is dissolved out. If the espanted amylic alcohol be distilled off, the resolutal morphine may be titrated, or the difficulty avoided by using magnessa unstead of ammonia.
- 3 A volumetric determination of morphine may be made by means of Mayer's solution, as described on page 140 The method has little practical utility

Further information on the determination of morphine will be found in the section on the assay of opium

4 Morphine readily reduces ferricyanides to ferrocyanides, with formation of pseudomorphine (oxydimorphine) —

4(2₃Π₃NO₃EG1+4K₃EoO₇=2(Ω₃H₃N₃O₂EG0)+8K₃EoO₇+H₃FoO₇. Consequently, on adding to the solution of a selt of morphine, slightly additated with hydrochlone scid, a mixture of aqueous solutions of ferric chloride and potassium ferrogrande, a blue coloniant on preceptiate of Prinsand blue is produced. This restino may be conveniently employed for detecting morphine in presence of the emchana bases.

- L K 1effer (Amad Chem Pharm, cm. 274) has proposed to tublase the reaction with ferrorpande for the quantitative determination of mosphine. For this purpose he adds a known weight of solid potessium ferrorpande to the morphine or its salt, and mixed them in a mortias with a minimum quentity of water. The contents of the mortiar are insed into a flask, potessium holide and hydrochlore said added, and the liberated volume determined
- ¹ There is some evidence that morphine forms a compound with amylic alcohol not decemposed by evaporation at the ordinary temperature (*Pharm. Jour.*, [3], xviii. 161)
- ² A solution of morphine in hydrochloric and cannot be shaken with amylic alcohol without axtisction of some of the alkaled, probably in the form of hydrochloride.

by deconormal codum throsubphate (hyposubphate). The difference between the volume required and that used in a blank apperment with the same weight of potassium forneyands corresponds to the salt roduced by the morphism. One ac of difference in the $\frac{N}{10}$ throsubphate used represents 0393 of anhyticous morphism's

Venturing (Gaz. Chim Ital, xvi 239) reports favourably of

Kieffer's process. The author's results were discouraging.

5. On mxxng a solution of morphune with one of solute dissolved in hydriothe and, a crystalline prequisitate is formed even in extremely dilute solutions. Under the microscope the crystalline form is characteristic of morphine, which may thus be distinguished from paparerin and codons, which bases along ore crystelline precipations with the reagent, while narcotine, narceme and thebame yield amorphous precipitates.

6. Addition of chlorine or bromine water, followed by ammonia, occasions in moderately concentrated solutions of morphine a brown colour or red coloration gradually changing

to brown

7. Morphine and its salts reduce notice and with liberation of odine. This reaction is also produced by albummoid and various other organic bodies, so that it is not absolute proof of the presence of morphin. The test becomes much improved and increased in doheacy by the following mode of operating —.

To the solution to be tested for morphia, as nearly neutral as possible, is added one of rodic acid in 15 parts of water. In presence of 1 part of morphia in 20,000 of liquid a yellow coloration is observed. In moderately strong solutions of morphine addition of starch-liquor gradually changes the yellow colour to blue, but not in solutions containing less than 1 per 1000 This is important, as with other reducing agents the blue colour is well marked in far more dilute liquids. On adding excess of ammonia to the yellow liquid the colour is discharged if due to foreign matter, but distinctly deepened if due to morphia If a solution of morphine, which is too dilute to give a blue colour with rodic and and starch, be mixed with these reagents, and some highly dilute ammonia allowed to flow from a pipette on to the surface of the liquid, two coloured rings make their appearance at the function of the fluids. A blue ring is seen in the lower acid layer and a brown one in the upper alkaline portion If a dilute solution of morphia be muxed with one of starch, and evaporated to dryness in a porcolain crucible at a gentle heat, and the residue, after cooling, he

¹ It is possible that Kieffer's process might be applied to the amylic alcohol solution of morphine, by agitating it with polassium ferricyanide solution. In such a case, summonia, if present, would not interfere.

moistened with rodic acid, a blue colour will be produced in presence of 1-20,000 of a grain of morphia (A Dupré).

Another way of employing the test is to agitate a solution of olde and with an equal measure of cathon disulphine, but an expanding the substitution of the sound of the substitution of the second colored even after adding a dop or two of didute implicit on the tested of morphine be now added to the mixture, and the whole again shake the carbon disulphined will be found after separation to have a world colour from dissolved colour from dissolved colour from dissolved colour from dissolved and metation of the amount Morphine can be recognised in this way in a single diop of paragone or tancture of opinion

Stein and others have described a colormetric method of estimating morphine, based in the rodic acid reaction

In employing the rodic and test it is essential that the resent should not give free rodine on treatment with a drop of dilute sulphune or acctic acid.

§ Solutions of morphine salts give no crystalline precipitate with either potassium chromate, thiocyanate (sulphocyanide) or ferroeyanide (distinction from strychnine)

APOMORPHINE, C17H17NO, When morphine or its hydrochloride is heated to 140°-150° C in a sealed tube, with a large excess of strong hydrochloric acid, or with zinc chloride at 110°, it is converted into the hydrochloride of anomorphine, the formula of which base differs from that of the parent alkaloid by the elements of water, though its formation is probably attended by polymerisation. Apomorphine may be obtained in a state of purity by dissolving the contents of the tube in water, adding excess of acid carbonate of sodium, and agitating with ether or chloroform, in either of which apomorphine is freely soluble (difference from morphine) The ethereal solution is separated and shaken with a very little strong hydrochloric acid, when crystals of the hydrochloride of apomorphine are deposited. These are separated, washed with a little cold water, and purified by recrystallisation. From its aqueous solution of the hydrochloride, sodium bicarbonate precipitates free apomorphine as a snow-white amorphous substance, readily soluble in alcohol, other, chloroform and benzene, which speedily turns green on exposure to the air The changed alkaloid is partially soluble in water and alcohol with emerald-green colour, in other with magnificent rose-purple, and in chloroform with fine violet tint. The colourless solutions of the unchanged substance soon acquire these tints. In its physiclogical effects, apomorphine differs from morphine in a very marked manner, being a prompt and non-irritant emetic. From 0 001

to 0 010 as the adult mediamal dose by the stomach Dangerous and even fatal symptoms have followed the hypoderume injection of 0012 gramme Apomorphine gives a crusson-red colour with funce acid, and brown with sold acid, but (cultake morphine) yields a nose-red or amethystime colour with ferric chloride, changing to violet and black. The most deluctar reaction of a pomorphine is the production of a green coloration when the solution is reacted faintly lakline with posssum hypotogen carbonate and exposed to the air. With a solution containing I part in 100,000, the green colora appears within ten munites.

Apomorphine is said to be liable to be formed in old solutions of morphine hydrochloride, which consequently acquire emetic properties; but the statement is disputed by Dott, and requires confirmation (Phan m Joses, 13], xvi 287, 299, 604, xvii 80).

Apomerphase Hydrochlould, O₂rH₂NO₅HCl, forms anhlydrous, munte, alming oystals, which turn greensh on exposure to light and an. It is freely soluble in water and alcohol, forming a neutral solution, which turns green on boiling or standing, and keeps better if very faintly and The freshly-male aqueous solution should be colouries, or nearly so It is generally held that if a 1 per cent solution be emerall-green, the sample should be rejected for methcal vars, but D D D ot (Plann Jow [3], xxi, 918) has pointed out that the coloration is so intense that very little soluted change is theoly indicated. Mortell found an old solution which had been exposed to light for three months to act quite effectively.\(^1\)

Basic Associates of Morphine.

As already stated, oputm contains a large number of lases, some of which are present in very minute amount, or are allogother absent from some samples. The names, formules, solubilities, and chief colour-reactions of these alkaloids have already been given (page 294 to 300), and morphine has been described at length (page 309). The following are additional facts respecting the less unportant bases of opition.

Conaurs, Caste, NO, melts at 136° when crystallised from benzae, and 121° when separated from alcohol or ether 11 forms large sx-suded prisms, which can be sublimed 11 dissolves moderately easily in hot water, giving an alkaline solution 12s silts, which are amoniphous, give precipitates with caustic alkalies and

¹ Morrell finds that a patient who is made violently ill by \(\frac{1}{2}\) grain of a pomorphine hydrochloride administered hypodermically, can take \(\frac{1}{2}\) grain thrice duly in the form of pills. Apomorphine acts as a powerful expectorant in cases of dinous bronchitis.

CODEINE 321

ammonia, soluble in excess of either reagent with nutric acid, codamine gives a dark green coloration with sulphuric acid, and in presence of a minute quantity of ferric chloride a greenish blue For other colour-reactions and solublities, see page 301 et sea

Course Coles. C.,H_mNO, or C.,H_mNO(OH) CCH, Thas base has the constitution of a morphise methyl-selor. The relation of codeuse to morphise and synthesis therefrom are described on page 167. Its theoretical relations and constitution have been recently further investigated by K norz (Ber. xxii 181, 1113) and S traup and Wiagmann (Montiech, x. 782). Codune occurs in opium in proportions ranging from 0.1 to 10 are cent. 1

Codeme crystallises from dry ether or earbon disulphide in small unhydrous prisms. From whet it is deposited in well-defined cotcheder are ortherhombre prisms containing 1 aqua and melting under bothing water to an oily luquid. Anhydrous codene meltis at 100°-126°, and solidrifes to a crystalline mass on cooling Codene is somewhat soluble in water, requiring 75 to 80 patts of colid water, or 17 at the boling-point. It is readily soluble in alcohol, either, amylic alcohol, chlioroform and benzene, but is almost insoluble in petceloum spirit (compare page 301). Codene is as soluble in ammona as in water, a fact utilised to separate it from morphino, but it is pracheally misoluble in excess of caustie potan or soda, and is precupitated by these respects from its queene solution, if not too chitus. Solutions of codene are optically active, the rotatory power being moch affected by the nature of the solvent, and the presence and proportion of free and In adolicities obtions, = 12° in additions, = 12°.

Codens has a bitter tasts, and resembles morphine in its physiological action. It is official in the Butish and several foreign Pharmacopoucas, and is chiefly employed to allay restlessness, cough, and other symptoms for which optim is generally presembed, and when the latter medicine is not idealated. In philass, it appears to prevent and oppease the ticking irritation of the cough, without draminging the digestion. It is an important remedy in diabetes,

¹ Codeno is usually nobated from opum by precipitating the squeene votroot by calcium, dionde, evaporating and cooling the filtrate, elessiolying the deposited crystals of the hydrochlorides in water, and prosquisting the marphine by amounts. From the filtrate, after consentation, the codesine can be recovered by treating by precipitating with caustic alkali, and purified by oxystalisation from either.

² The hydroxyl-group in the codeme molecule does not appear to be phenoine, as evidenced by the insolubility of the alkaloid in caustic alkalies, and its negative reaction with ferro chloude.

and is also recommended as an hypnolic in mental disease. The official dose is from \$\frac{1}{4}\$ to 2 grams In larger quantities, codeme produces narcotism, often preceded by vormiting and occasionally by purgung.

Codome is a strong base, having a marked alkaline reaction, and forming crystallisable, soluble salts, which are neutral to litinus and methyl-orange The free base precipitates solutions of lead, iron,

copper, and certain other of the heavy metals

Colone High schlorade crystalines us raduated groups of prisms containing Birlick+ $2\Pi_{c,0}$ valuable in about 20 parts of cold water. The solution is law-rotatory ($S_{j}=-108^{\circ}$). The crystals lose a point of their water (j agua) readily, but the remainder is only driven off by many darys heating at 100° (S_{i} has id, j harmony j but easily at 120° (j bot). Hence the proportion of water in commencial samples of the salt is vanishly

Codeme Phosphates The sait BH_PO_4+2H_O is obtained as a crystaline presquate by adding ordenic to a solution of phosphane and till the reaction is only family acul, and then adding excess of alcohol. When recystallined from what the composition is unchanged, but the sait deposited from the solution in bot dilute alcohol contains BBH_PO_4+H_O. Both form loss their water at 100°, and are new with in commerce, as also a preparation containing excess of phosphore and. The usual composition of commercal redeemed phosphate is B_HB_PO_4+H_O. (Do t.) If the sait turn gray or yellow at 100°, the presence of impurity is in listed. The phosphate is said to be the preferable form of employing colores for Nyrodernes negocious

Detection and Determination of Coderne

In its reactions and general characters codeme presents a strong resemblance to morphine, but is sharply distinguished by its ready solubility in ether and chloroform, and its precipitation by excess of caustic alkali. Codeine does not reduce iodic acid, and gives no coloration with ferric chloride. In strong nitric acid it dissolves to a yellow liquid which should not become red (difference from and absence of morphine). With pure sulphuic acid, codeine gives no coloration, but on warming, or very prolonged standing (several days) at the ordinary temperature, a blue colour is developed. This colour is produced if a trace of nitrie acid, ferric chloride, or other oxidising agent be present, an assemate being the preferable reagent The blue coloration on warming with sulphuric acid and ferric chloride is apparently common to all ethers of the codeine class. Fronde's reagent (page 147) is stated by some observers to produce a dirty green colour, soon becoming deep blue, and changing in twenty-four hours to yellow, according to others, a cherry-red tint changing to roles, se produced L. Ra by estates that if solid codeme be sturred up with two diops of a solution of solution hypochlorits, four drops of stong sulpinare and added, and the whole mixed together, a splendul and persistent blue coloration results. Esculin was the only other substance (of thiny examined) which gave at all a similar reaction. La fon uses a solution of I gramme or amnonium selente in 20 cc of stong sulpinare acid, which gives a magnificent green colour with traces of codeme. Other reactions are green on person 30 to 30 cm.

Commercial codeline has been met with adulterated with ammonum tartiate (Pharm Jour, [3], xiv. 1035), which salt closely resembles it, but is distinguished from codeline by its insolubility in alcohol

Cl as as on has based a method of determining codene on the well-known fact that it completely decomposes morphine sells (A' Y Pharm Randschau, 1890, 40, Jone Chem. Sec., Ivini 1198). The warm aqueous solution of the free base is treated with excess of morphine sulphate with frequent shaking, and allowed to stand in the cold for it least twenty-four hours, when the disposited morphine is filtered off, dract, and weighed (or tribated). The amount fourst, multiplied by 0.9868, represents the sulpivious codeme, or by 1.0412, the hydrated colonic (C_hH_AC_h+H_AO). To separate morphine and codeme, the mixed bases, or their salts, are evaporated to dry-ness with excess of magnesses. The resulte treated with water, and the liquid shaken repeatedly with ether free from alcohol, the effect of the colonic off, and the residue exhausted with hot water. In the issultant solution the coddine can be determined as above described.

Classem (loc cit) has also pointed out that fice codeine conpletaly decomposes summonium salts when heated with them, and has based on the fact a method of determining the alkaloid, but as morphine behaves in a similar manner, the fact has little practical value.

The sumplest means of determining codeme and morphine in admixture is to preceptate the solution of the hydrochloudes with and carbonate of sodium, and wash the dred preseptate with chloroform The resultse counsate of morphine The aqueous filtrate is treated with causies sody, aguitated several times with chloroform, the varaous chloroform, used and extacts united, evaporated, and the resultad codeme dried at 110°, and weighted (D. B D ot 1).

Pseudocodeine, C₁₈H₂₁NO₃+H₂O, was discovered by E Merck in preparing apocodeine (Arch Pharm, ecxxix 161). It is a strong base, crystallising in needles melting at 178°-180° It is lavo-rotatory, forms crystallisable salts, gives no reaction with ferric chloride, and has a physiological action similar to, but weaker than, that of codeins

Appendame, C₂H₄NO₀, as saul to be produced by heating codume hydrochlorate with a concentrated solution of zan chlorate for fifteen manuter. It is described as gammy, insoluble in vater, a soluble in alcohol and either, and yndding amorphous salts in hyavological scision it is a valuable expectional and mild emetach. Appendeme gives a characteristic blood-red colour with intra colour. In B D of the dubt is the existence of appendicume, and states that commercial opendesse hydrochlorade is not of a vory definition in the commercial opendesse hydrochlorade is not of a vory definition in the commercial opendesse hydrochlorade is not of a vory definition of ordenia, polymersed bises, chlococolide, and apomorphics in Palysnological results appear to harmonies with this view (Pharm. 1998; 13) XX 1878, 918, 956, 996)

Methocolesies or Dunethylmorphine, C₁₇H₁₇NO(OCH₂)₂, is of interest merely from its theoretical relation to morphine, codeme and thebaine (compare page 296). It is a base forming hard brilliant lamme melting at 119°, and yields with sulphinic acid a

brown coloration, turning violet on addition of water

CRITOPINE, C_BIL_BNO_B occars in but very small quentity in opium, and is pecepitated on adding causte sools to the mother-liquor from which codeune, naiceme, tholeane and papaverne have been separated. It crystallises from alcohol in minute six-aded passes. It is optically inactive, sparnely soluble in boiling alcohol, very sightly in benzene or petroleum spirit, but more readily in oblocoform. When freshly inscipated it is soluble in their, but slowly separates from the solution (See also pages 301, 504) Cryptopus and its salls have a butter taste, and pungent cooling affect-state; they are hypnotics and mydratics

Cryptopine salts when dissolved in hot water usually produce on coulding a gelatinous mass, which is gradually changed to crystals. The normal subjects does not crystaline; the acts sait gelatinues, as the solution cools, and the jelly shows but shight signs of crystalinus, even after standing swerall weeks. The acts contains and cast desirate are very sparnely soluble. Neutral cryptopine nucontes, (Ca₁H₂NO₂N_CH₁O₂+1OH₂O, is mobible an coid, and but slightly soluble in boiling water, and is probably the form in which the alkaloid exists an opinum (Plans 10ar, 13 X vizu 250)

DEUTEROPINE, CookerNO an alleged homologue of protopine

and oryptopine, requires further examination.

GNOSCOPINE, Ci₂₄H₂₆N₂O₁₁, occurs in the mother-liquors of naiceine When recrystallised from boiling spirit the base forms long, thin, white needles, having a woolly appearance when dried. It melts at 233°, decomposing at the same time, and burns with a manky flame, leaving a shelden of cherood. In pure sulphure and, gnoscopine dissolves with slightly yellow colour, which becomes at once armine-red upon addition of a time of nature acid, the colour being permanent. This section distinguishes the base from rhoaddine, which becomes red with sulphune or hydrochlorus and alone (Pann Non. [3], ix 82). Gnoscopine hydrochlorus gives a buff-coloured precepitate with platinic chloride. (See also noze 301)

HYDROCOTARNINE, C₁₂H₁₈NO₂, is formed from narcotine, together with meconin, by the action of nascent hydrogen. It volatilises partly unchanged at 100°, and forms readily soluble salts

Linymopris, C₂₈H₂₈NO₆, is obtained from the mother-liquous left from the preparation of monthine by the Robertson-Gregory process (see page 308). It is a weak base forming no acelate. It is coloured connege-red by mitre sauk, and pale violet by subjunct and, the latter colour changing to a dark brown on heating (See also naces 301, 304).

LAUDANINE, Con HornOs, occurs with lanthopine It has recently been prepared on a commercial scale by Merck from onium mother-houors, but the yield is only one-third that of ervotopine Laudanine crystallises from its solution in boiling alcohol in transparent granules or hexagonal prisms melting at 166°. Laudanine is levo-rotatory, tasteless, and poisonous, the hydrochloride being bitter and resembling strychnine in its effects. It resembles morphine in dissolving in caustic alkali solutions, but the sodiumderivative is remecipitated in glistening white needles on adding excess of caustic alkalı From its solution in caustic alkalı laudanine is wholly unremoved by chloroform or amylic alcohol, but is extracted if precipitated by ammonia. Its phenolic character is further evidenced by the green coloration yielded with ferno chloride. Treatment with methyl rodide converts laudanine into a base chemically resembling codeme, and distinct from laudanoane. The solution of laudanine in pure concentrated sulphuric acid has only a very faint pink tint, the same acid containing iron vields a slightly deeper tint, but on heating either solution till the acid begins to volatilise, a violet coloration is obtained. With mitric acid, laudanine gives an orange-red colour Laudanine is a strong base, having an alkaline reaction, and forms well-crystalheed ealts of a bitter taste BHI is sparingly soluble in cold water, and BHCl easily soluble in water, but nearly insoluble in brine. (See also pages 301, 304)

LAGDANOSINE, C21H27NO, 18 homologous with laudanine, but 18 not produced by heating that base with methyl iodide Laudano-

sina is isolated by convenion into its sparingly soluble hydraditie. It crystallness from benzene in needles melting at 91° Both the free alkaloid and its salts taste very bitter, and are tetanic poisons. Landanesme is dextro-rotatory. The solution is strongly alkaline. It gives no coloration with fearne chloride. (See also pages 301, 304). Morenius, $C_{17}H_{10}NO_{27}$ has already been fully described (page

309)

Miscontines, C., H., NO. (1969 301), forms a brownish yellow amouphous mess, soluble with difficulty maintonia, but readily a caustic alkales. The base cannot be removed from its solution in caustic solar by agricultura with ether, but is extended from its ammoniacal and line-water solutions. Mecondulue is alkaline in reaction, and menty destitute of tiste, but yields very hitter, unstable salls. It is very existly decomposed by unicard acadily, but production of a rese coloution. It is alsosived by strong subplume each with an observement and but untreased with the otherwement and but uttree only with an orange-red colour.

NARCHINE C2, H20NO2, OF C12H20NO4 GO C6H2(OCH2)2 COOH (compare page 299) This base was originally discovered by Polletier, who attributed to it the melling-point 92°C, but Hesse found it to melt at 145° This latter figure, although subsequently corrected by Hesse himself, has been generally adopted by compilers, though Claus and Merxner found 162°, but E. Merek has shown (Chem. Zest. 1889, p. 525) that the ordinary commercial alkaloid of English manufacture melts between 150° and 160°, and the pure base at 170°-171°. Narceme crystallises from water in long white prisms or delicate needles, containing 2H₀O₄ which is driven off at 100° It has a bitter taste, with styptic after-taste, and powerful hypnotic properties. It is optically mactive It is very sparingly soluble in cold water or spirit, but dissolves very easily on heating. It is but slightly soluble in chloroform, and insoluble in other and benzene Narceine is precipitated. on adding ammonia or caustic potash to solutions of its salts, but dissolves in excess of either reagent, and on addition of a large excess of caustic alkalı is reprecipitated as an oily liquid 2

Narcence as a very weak base, the free alkaloid having a very feeble alkaline reaction to deheate litmus, the solutions of its salts may be titiated with litmus just as if the alkaloid were absent. The accitate is decomposed by water, and the base is said to be extraordal by chlorofoum (but not by amyle alcohol) from hyduid contaming

¹ Dott states that the melting-point is indefinite, as partial decomposition

² Nauceine containing a carboxyl-group, its solubility in alkalies is normal, but it seems probable that the oil precipitated by excess of caustic alkali is an alkaline are or in at a ratio r than the free alkaloid.

even free mineral scule IHCI forms needles or short stout prisms very easily soluble in water and alcohol, and melting prisms revy easily soluble in water and alcohol, and melting the decomposition at 163° Nareane blorated from the hydrochloride or other salts by ammonia rotains hydrochloric soid with great persistency, and cannot be purified by recrystallization from water or dilute alcohol According to E. Merck (Olem Ezt, 1889, p. 525, Plarm Jour, [3], six 1034, xx 481) narcane can best be obtained pure by crystallization from water containing some ammonia or canste alkelli, but a considerable quantity remains in permanent solution. For theirspettic purposes, the presence of a small proportion of hydrochloride is of no consequence, and Merck considers that a preparation froe from meconin, and so far freed from base sall as not to much below 165°, as sufficiently guita.

Chlorme-water, followed by ammonus, grees a blood-sel colour with naccens, but many other substances (e.g., tamm) lebax a smilarly Potassum bichronate grees a crystalline precipitate after some time Toline grees a hown paccipitate in nationic solutions, but if ammona be added to remove excess of notime the pracquisties are seen to be blue Week rodine solution colours nan-cense black-blue, in boiling water a colourless solution is obtained, but the crystalts formed on colouing have a violet on blue colour Sulphure acid containing took and grees with narcenne a black coloration changing to red (see also sugge 30.9 of see)

"Me con a reina, as "according to E. Me re k, as a preparation of a very variable character, of which on a form consists of a yell-lowah liquid containing codems, narceine, and an undenfated and soluble in ether, but no mecoure acul. In another case the "meconicone" formed a white powder melting at 110°, and consider, and a distinguishment of a mechanical mixture of narceine and meconic sond, which on adding water combine clientally, and the inerpstallized products melt with evolution of gas at 126°, which is the melting-point of acid naiceine meconite (Plan m. Zed. 1889, p. 90)

Margories, C.3H_3/O₂, occurs in optim in very variable quantity, the usual range being from 1.3 to nearly 11 per cent, but some samples contain these too minute to be recognised by the usual methods Natootine may be extracted from dread optim by ether or beneare, or by the same solvents from the precepitate produced by aumonia in the aqueous solution of optima. It may be separated from nateous by precipitating the solution with excess of aumonia, when the navoeme remains in solution.

Narcotine crystallises from alcohol or other in colourless, transparent, glittering prisms or groups of needles, which melt at 170° ,

Optum from which the narcotine has been removed in this manual is now an article of commence.

and resolidity at 130°, crystallising if cooled slowly — Above 200° in narcotine is decomposed into in eo on in and octarinis 1° life fieldly navootic, exhibiting possonous effects only in somewhat large doese (1 for 50° grainnes). The solid base is nearly tasticable, but the solutions are latter. In the free state narcotine is invortatory, but the salts arithat destro-rotatory.

D B D ot t has obtained the scattats, sulphate and hydrochlorate of nacotine in a crystalline state; but the first of these saits a success completely decomposed by solution, the base being precipitated and free acetic sent formed. The same reaction occurs when solution scattate is added to a solution of introchloride of microtine hydrochloride (compare page 306). The hydrochloride and sulphate of introches as somewhat more stable, their solutions remaining client even when largely dulated, but they react with himsu just as if the and were uncombined, and yield the nacotine to chloridom and similar solvents. These facts prove the basic properties of nacotine to be very feebly marked.

Narcotine meconate forms a syrupy solution, which on evaporation dries to a varnish which redissolves perfectly in water

The caustic alkales, alkali-metal calbonates, and ammons throw down narcotine as a white crystaline precipites, almost insoluble in cold wates and in excess of the precipitants. It may be extracted from the alkaline highed by chloroform to heimen, or less readily by either or amyhe slochol. It is practically unaffected by petroleum spirit (compare page 301).

Narrotine is precipitated by the usual alkaloidal reagents, but the reactions are not very chanacteristic. With potassium thiosynate it it yields a crystalline procipitate readily soluble in acids, even in acetic acid. Iodised potassium rodule precipitates naicotine from extremely dilute solutions. Narrotine may be precipitated and intractly by Mayer's solution (page 139).

If a solution of narcotine in dilute hydrochloric acid be treated with bromine, a yellow precipitate is obtained, which dissolves on boiling, by gradually adding bromine-water, and boiling, a fine rose

2 Hesse found for the free alkaloid-

			Alcohol	Chloroform and Alcohol	Chloroform	
	Concentration,	٠	074	2	2 and 5	
	Sp, .	٠	-186* 0	-181° 5	-207 8	

For a solution in "benzine" Dott and Preddie found $S_0=-229^\circ$ (when δ was 1 5), and for a solution in dilute exalic said, $S_0=+62^\circ$

¹ The constitution and decomposition-products of narcotine are described on page 298

Surroutine hydrochloride is neutral to methyl-orange (D o t t).

colour is produced, but is readily destroyed by excess of bromime. The reaction is characteristic With chlorine-water, narcotine gives a yellowish green colour, turned orange by ammonia. Iodic acid gives no coloration with narcotine. If narcotine be mixed with twice its weight of cane-sugar, and the mixture moistened with strong sulphure acid, a fine and persistent mahogany-brown coloration is produced, said by M Robin to be highly characteristic. (See also page 302)

OPIANINE, to which the formula Co. Ho. NO. 18 attibuted, 18

probably merely impure naicotine

OXYNARCOTINE, Coo Hop NO as a contained in the mother-liquois of narcotine.1 It forms minute crystals, somewhat soluble in hot water, but little soluble in hot alcohol, and insoluble in ether, chloroform or benzene. By exidation with feiric chloride it yields cotarnine and hemipinic acid BHCl+2H₂O forms

crystals (See also page 101)

PAPAVERINE, C20H21NO, is a weak base of feeble narcotic properties It is separated from narcotino by crystallisation from a strong solution in exalic acid, the acid exalate of papaveine being very sparingly soluble Papaverine crystallises in thombic prisms or needles, or sometimes in scales. It is slightly levo-rotatory,2 though its hydrochloride is mactive. The neutral succinate forms large tabular crystals melting at 171°, and soluble in hot water, the benzoate, triclinic crystals melting at 145°, and soluble in alcohol but msoluble in water, and the saliculate, monoclinic crystals melting at 130°. Sulphume and containing rodic and gives with papaverme a purple colour, turning black and green. Dilute solutions of papaverme salts are not precipitated by phosphomolybdic acid. Tincture of rodine, added to an alcoholic solution of papaverine, gives gradually a precipitate of crystalline needles. With potassio-iodide of cadmium, papaveime vields a dense white precipitate (See also page 301 et sey)

Papaverosine, found by Deschamps (1864) in the dried seed capsules of the poppy, crystallised in prisms, was soluble in alcohol. ether, chloroform and benzene, and formed a gummy hydrochloride

With sulphuric acid it gave a violet coloration

1 Oxynarcotine was first isolated in an impute condition by D. Biown. from orude narcome. This product was purified and analysed by Alder Wright and Beckett.

² G Goldschmidt (Monatsch, ix 42) states that pure papaverine is mactive, and suggests that the optical activity of landamic should be remyestigated, as these two alkaloids constitute the only two known exceptions to the Bel-Van't Hoff theory that derivatives of optically active substances are also active.

PORTHUMENTA, described by Mersk in 1887 as the red colouring matter of opuns, according to Hesse as a mutatre of several bases, one of which is meconidine, and another probably via ordine, which latter alkalond also occurs in the oppsiles and other parts of the red poppy. Kanny Lall Dey (Pharm Jowes 1987) as 1997, states that by treating the enqueue extract of Indan opuns with aumonia or sodium carbonate, and immediately against my with either, the ethereal solution always leaves on evaporation a body (rhossbire!) which, when warmed with dutue hydrochione and, given a ruch purple coloration, and he recommends the reaction as a test for Indian opuns. With Turkey and Smyrna opuns no such incetton is obtained.

Theorems, $C_{00}M_{11}NC_{00}$ appears to be the most widely-distributed of all the optima likeholise. It is formed in very immisste quantity in optim, but has been met with also in Madega condata, Siftophousin alphyllian, Sangulana a Canadersis, and Cheldonium angus. Protopine tesembles varylopine, but the solutions of tesalts have a bitter taste, and do not gelitimise on cooling. In small doses, protopine acts in fogs as a narrowise, and in stronger doses paralyses the muscle-substance, and the pruphetal ends of the nerves. Upon mammals it has a porsonious setton like that of complion, but differs from it in paialy sing the enrediating organs (See also pages 301, 304)

Pestrovonetrix Oxydmonphme C_hH₂₀N_cO_e* The akteloat is best puriod by solution in ammoun, from which it crystallises in coloniless cruets or delicate silky needles containing 3 agas. It as very weak bees, forming no accitate, and as without action on vigetable colonis. It is taskeless and not posonoms. It dissolves readily in cauri-to alkalies and milk of Inus, but is involutible in all the ordinary alcoholic and ethereal solvents, as also in dicite sulphine sent and alkaline carbonates. (Compare page 301). Its most soluble sails its the hydrodilorids, which requires 70 posts of old water for solution. On adding ammona, avoiding excess, the alkalor dispersible the art great the two propositions are presented in a greating set from the lot, and in a gelatimous state from the cold solution. He is a finish that when pseudomorphine is mixed with an equal weight of canceraga, and

Merck repeatedly dips a slip of filter-paper in the ethereal salution, alioning it to dry spontaneously after such immersion. The paper is then moistened with hydrochlorio acid and exposed to steam, when it will acquire, especially after drying, a more or less distinct reso-red colour.

² Pseudomorphine occurs very rately, having been observed by Hesse in good Smyrna opinim only once in four years. It may be prepared by treating morphine with oxidising agents of moderate power, such as potassium ferricyanule or dilute permanganate (page 144).

strong sulphurne acid (pure) added, a characteristic dark green coloration is obtained, which gradually turns brown (compare test 5, page 315) If the acid contain a minute quantity of iron, a blue coloration changing to green is produced

RHEADINE, Col Houndon exists in all parts of the red poppy (Papaver Rhaas), and in the ripe seed-capsules of the white poppy. It forms small white prisms, which are testeless and not poisonous, Its solutions in weak acids, avoiding excess, are colourless, but on adding excess of sulphuric or strong hydrochloric and a purple-red colour is produced. This is destroyed by alkalics and restored by acids, and is so intense that I part of rhosadine will colour 10,000 parts of water purple-red, 200,000 deep rose-red, and 800,000 distinctly red, although only a fraction of the base is converted into colouring matter. The colourless solution of rheadine in acids is precipit ited by tannin. On adding potissium iodide to a solution of the acctate, the hydrodide is precipitated as a dense crystalline mass, consisting of microscopic prisms. An aqueous solution of theadine becomes red by prolonged boiling, part of the alkaloid being converted into the isomeric base in cagenine (soluble without colour in acids), and on adding a drop of hydrochloric or sulphung acid the whole base is decomposed, the solution acquiring a purple-red colour Cold dilute sulphuric acid converts solid theadine into a colourless resmous mass, which soon dissolves with splendid purple colour, changing to dark purple on boiling, and depositing on cooling small prisms which are brownish red by transmitted and green by reflected light, while the limit retains rhosagenine equal to 99 per cent, of the rhosadine present, together with the colouring matter

Opum sometimes contains a base which gives the above colourreactions with sulphunc acre, but it is somewhat doubtful if it is actually rhoadine (Compare Porphyroxine, page 330)

actually thoseline (Compass Porphyroxme, page 330) Thumanra, $C_{10}H_{11}NO_{2}$ or $C_{11}H_{12}NO(0$ $C\Pi_{12})$. Thebanne occurs in opinin in proportions ranging from 0.15 to 1.0 per cent. It crystallises in silvery seeles from dilute alcohol, and in needles or hard quadrate prisus from stong alcohol. Thebanne melts at 193°, and is not sublimable 1.1t has a sharp and stypic testle, and is a powerful tetaine posson, producing symptoms resonant bloos due to staychinue. The fatal does is smaller than that of morphine. Thebanne gives a reddles hown coloration with chlorus-

¹ This is Hesse's experience, and is confirmed by Dott According to other obeatous, at about 180° it sublimes without fusing, and is deposited in minute crystals revembling cafform; while at higher temperatures, needles, cubes, and prisms are obtained.

water and ammonia. Its other colour-reactions (and its solubilities) have already been described (See page 301 et seq.)

Thekame is stated to be extracted (with some difficulty) by-chlorform from its east solutions, but the statement requires confirmation, as it is inconsistent with the strongly-merked bene characters of thekame 1 From mercoine, thekame may be separated by treating the concentrated seeine solution with excess of barn lead scattes, which preceptates the narrotine only. Dittie scatts readily aller thekame, conventing it into the isometric bases to be nine and the bair of the strongly solution in heal alcohol and moscitate in other sample solvents. When heated to 90°, under pressure, with funning hydrochloric acid, thekame resides a base having the probable formals Ch₂H₁N₂O(OH)₂, colled by its discovers, W C. H o wait of (Ber, xvii 527, xvi 1599) in or phothe bairs, by undeast test cognian and relation to morphism

TRINGERS, $C_{\rm eff} H_{\rm e} N_{\rm e} C_{\rm h}$, was soluted by Kauder m musticularity from the nother-lequence of the opun-alkaload manufacture. It resembles morphine and laudanine in being soluble in soda solution, but is representated in the form of an oil by a large screen of the reagent. Tringene crystallies in characteristanhydrous, transparent, needle-like plates incling at 182°, easily soluble in chloroform, but only sightly in ether With sighility soluble in chloroform, but only sightly in their With sighility soluble in chloroform, but only sightly in the FW with sightly soluble in chloroform, but only sightly in the With sightly soluble in the control of the sightly sightly in the With sightly in the With sightly soluble in the control of the sightly sightly in the With sightly sightly sightly sightly in the With sightly sightly sightly in the With sightly sightly sightly in the With sightly sig

Opium.

Opum is a gummy mass, consisting of the inspisated juice from the incised unitpe fruit-capsules of Papaver sommiferum, hardened in the air

Opmus is produced in Turkoy, Ana Minor, Pessa, Jinda, China, and other countries, but Sunyan, Constantinople, or Turkoy opnus is the only variety recognised by the majority of the phanmacopcans. Persian and East Indian opnuss are imported chiefly as sources of the opnum alkaloids. Chinese opnus is wholly consumed locally.

1 It is possible that certain thebains salts are soluble in chloroform (as all those of codeme), and also dissolved as such by agitating their aqueous solutions with chloroform

³ The vanety of poppy cultivated in Ann Mino is said to be the Mond, within usually has pumple flowers, and black, though continually white, needs It is said to be usually riches in morphis than that from the white-flowramp and white seeded poppy, which is nich in manodane, and appears to be the only kind cultivated in Seprit, Feren, India, China, and Japan. (For a chemical dustratedon between Tunley and Indian opum, see page 350.)

Opium varies considerably in appearance, composition, and quality, according to its origin and mode of preparation 1

Opum is remarkable for the large number of definite, highly complex, cyraldine principles contained in at. Of these the majority are at kaloids, a list of which is given on page 204. In addition, opum contains a cetia, Isctic, and me conic as oil, as he last substance being pecular to opum. Besules these bodies and the inorganic constituents, opum also contains the indifferent bodies me conin, me con oil osin, and opion in, and a variety of sugar, together with guinny and pectous matters, albumm, wax, fat, doutchous, resun, and a lumond sach. Woody fibre and other extraneous matters are also frequently present; but genume onum is wholly free from both starch and tame.

The following may be taken as the general composition of opium:-

Per Cent. 1	Per Cent
Morphino, 8 to 15, average 8 Gun Other elikaloids, 9 5 to 2 Store Inso	a and soluble humoid 40 to 50

ALKALOUES. Morphuse is the most sbundant of the bases of opium, and the most valuable of the constituents. Most of the pharmacopease require dried opium to contain not less than 10 per cent of anophine. Good Surpras opium deprived of vaster usually contains from 12 to 15 per cent of morphine, though cakes from the same case are apt to vary considerably, but if the proportion be below 10 per cent on the dry substance, adulteration may be suspected Expythan opium is poorer in morphine then that from Asia Minor, this proportion ranging from 6 to 12 per cent, but it contains a larger proportion of narrotine Persans opium is extiemely variable in quality, probably partly in consequence of this protice of mixing it with sagar and other saddlerants, though much of it is equal to ordinary Turksh opium. East Indian opium is as a tule, ranktably weak in morphine, the proportion beautre.

The product of Assa Mison is described in the Bratisk Planranospana (1885) in a follows —"In rounded, irruptally formed, on Historian dissease, reprincipled to weight, but commonly about eight ounces to two pounds, usually forward with protices of Poppy Jeaves, and seathered never with the redshist-how on daily fruits of a species of Russex. When fiesh, pleasts and internally somewhat mostly counsely granular, and redshists—one electrical-toward, but becoming harderly lesspung, and darkening to blackuh-known. Odour strong, peculian, nanotic, teste nanisonally britter."

sometimes as low as 24 per cent, more commonly between 34 and 5, and ceassonally as high as 8 or 9 per cent. This inferencity is probably partly due to climate and partly to defeative methods of collection and preparation. The variety known as "Patina garden opium" is prepared specially for method use, and contains from 7 to 8 per cent of morphine. In Chinese opium, the proportion of morphine is generally low. Finath opium valided Gui bourt from 144 to 22 s of morphine, and German from 165 to 20 per cent; that from the white poppy containing, according 16 1111, 6 f per cent (f) Algernan opium from red poppies yielded 104 to 178 per cent (f) In United States opium, the proportions of morphine and from white poppies 15 to 8 f per cent (f) In United States opium, the proportions of morphine cohervol have ranged from 7 4 to 10 2 per conts.

The morphine in opinin is usually stated to exist in combination with meconic acid, but Dott has shown that morphine ordinarily exists in opinin partly as meconate and partly as sulphate ² In some cases taccos of acetate and lactate are present

Naroctine exists in opium in widely varying proportions and often in considerable abundance. Upwards of 10 per cent has been occasionally met with. East Indian opium always contains more naroctine than morphine, whilst French opium sometimes affords neither naroctine, narocine, nor the lesium.

The narcohne in opium is genorally assumed to be uncombined, as it is readily extincted by tocating the original (dreat) substances with other or benzene, but as narcotine is readily removed from the cardilated solutions of most of its salts by agitation with a custable solvent, such as chloroform or benzene, it does not follow that its extraction from opium is due to its presence in a fee state. It most probably ausily exists as meconate Oossimolity the narcotine resists the action of solvents, unless the sample of opium has been pureyously treated with ammonia.

¹ Aubergier states that in one case the product contained 13 per cent of morphine, while the opum from a neighbouring farm, where the collection was made somewhat laker, contained only 11 per cent

8 Twolve samples of opuum analysed by Fluckiger (Pharm Jour [3],

Positypycoine, according to Kanny Lall Dey (Pharm Jour., [3], xxi 397), is a definite basic substance, always pissent in Indian opium, but absent from Turkey or Smyrns opium. Dey regards its presence as so constant and chanceleustic of Indian opium that he utilises it in toxicological investigations (See near 330).

The other alkaloids of opium have been observed in the following proportions —

Meconin is an indifferent body, crystallising in colourless, shining, six-sided prisms, which melt under water at 77° C, or alone at 110°, and distil at 156° It is odouless, bitter, and readily soluble in alcohol and chloroform, but only sparingly in other

v 845) gave the following analytical results. The proportions of morphine are most probably sensibly below the truth

Description of Optum	Ethereal Ext	ract, consisting of	Pure	Morphine	
Description of Optum	Wax	Crude Narcotine	Natcotine	Crisde	Pure
1 Paine, 2. Indian (1852-59), 3 Akbari, 4 Behan, 5 Behan, 7 Hydernhad, 8 Caudeish, 9 Perstan, 10 Egyptin, 11 Playford Suffolk (1898), 12 Soguith (1869),	14-2 12-7 18-5 13-6 6-5 0-4 16-7 14-8 11-5 8-8 12-0	10 0 0 0 8 8 5 7 6 7 6 9 7 10 2 12 2 9 8	40 055 457 811 547 647 681	11-2 11-2 14-2 10-6 14-1	8 0 4 3 8 6 6 1 8 8 8 3 7 1 5 8 4 8 8 8

assays of thuty-eight samples of opium, published by M. Ad it a. n., showed a proportion of morphise exceeding 7 pe nor in all but two schemes are specified to the contract of the proportion of morphise and the sample showing only 3 87 per cent of morphise contained 3 8 of narcotine, while other samples contained on 10 per cent of morphise and only the same precented or narcotine. This variation is doubtless this ensous why some samples of opium cause little or no headeden and other concession very dispersionly symptoms.

Narceine often occure more abundantly than thebaine

Meconin may be readily crystallised from boiling water, in which it is moderately soluble

The meconin contained in opium, in which it exists in the proportion of less than 1 per cent, is probably a decomposition-product of narcotine, from which base it may be prepared by heating with nitric acid.

Meconin is extracted from its acidulated aqueous solution by agitation with benzene, chloroform, or amvhe alcohol, the firstnamed solvent being preferable. Meconin dissolves in concentrated sulphuric acid, without at first producing any coloration, but the solution gradually assumes a greenish tint, changing to reddish in the course of twenty-four hours. If the hand he then warmed. the colour changes to emerald-green, blue, and purple, finally becoming red. The shades and order of the colours obtained depend much on the proportion of acid used, the tints being bluer and the reaction more delicate with a small quantity. Evaporated with slightly diluted sulphuric acid, meconin gives a gicen colora-In concentrated hydrochloric acid it dissolves without change of colour, even on heating. If meconin he dissolved in strong sulphuric acid and a minute fragment of potassium nitrate added, a vellow coloration is obtained, lamidly changing to a fine scarlet, which fades slowly and is changed to vellow on heating, The reaction is delicate

An aqueous solution of meconin gives precipitates of characteristic microscopic appearance with rodused potassium rodule and a solution of bromme in hydrobromic acid (T. G. Wormley)

MEGONOSIN, C., H., D., was obtained in brown, leaf-like crystal-ine masses from the mother-liquous left on the molation of meeconin. When pure it is coloulies, freely soluble in alcohol, ether, and hot water, fuses at 88°, and on evaporation with somewhat diluted sulphure scale yields a red colour, changing to purple

Orionus, according to Hease, as contained in small quantities in Smyrne opium. It founs white needles which nelt at 227° and contain no introgen. It is insoluble in water, but dissolves in alkalies, alcohol, and either. When boiled with milk of lime, opioum is decomposed, an acid being formed which is friely slotled in water and ethor, and gives a bulky precipitate with lead acetate in alkaline solutions.

Μεσονίο Ασιρ, $C_7H_4O_7=C_8HO_2(OH)$. (CO $OH)_2$ This substance is characteristic of optum, in which it exists cliefly in combination with the alkaloids, but sometimes a portion of it appears to be present in a free state.

Mecome and may be prepared from opium by precipitating the neutralised aqueous solution of the drug with calcium chloride. filtering, and decomposing the mecunitate of calcium meconate by repeated treatment with warm diluted hydrochloric acid. A preferable plan is to precipitate the aqueous solution of opium with neutral lead agetate, filter, suspend the precipitate in water, and decompose it with a stream of sulphwetted hydrogen. The filtered and concentrated solution deposits meconic acid on addition of hydrochloric acid. The product may be purified by re-solution in hot water, cooling, and adding hydrochloric acid. Meconic acid may also be conveniently prepared by precipitating it as the calcium salt, decomposing this with a slight excess of oxalic acid, filtering, and concentrating

Meconic acid crystallises in micaceous scales or small rhombic prisms containing 3 aqua. On being heated to 100°, it loses its water of crystallisation and leaves a white effloresced mass. At 120° C. it splits up into carbon dioxide and comenic acid. C.H.O., which at a higher temperature again loses carbon dioxide. and forms pyromeconic acid, CoH, Og 1 Comenic acid is but sparingly soluble in hot, and is almost insoluble in cold water. In absolute alcohol it is quite insoluble Meconic acid dissolves in 115 parts of cold, or 4 parts of boiling water, its solubility in the cold is diminished by addition of hydrochloric acid, which therefore causes a precipitate in strong solutions. When the solution of meconic acid is boiled for some time, especially if hydrochloric acid be present, comenic acid is formed, and civstallises out as the liquid cools The aqueous solution of meconic acid has a sour astringent taste, and strongly acid reaction

Meconic acid is freely soluble in alcohol (distinction from comenic acid) and is deposited in fine crystals on spontaneous evaporation of the solution It is much less readily soluble in ether and is almost wholly insoluble in chloroform

Nitric acid readily acts on meconic acid, much oxalic acid being formed

Meconic acid derives its chief analytical interest from the fact that it is strictly peculiar to opium and its preparations, and hence

Comenic and forms prisms, lamine or granules, insoluble in alcohol. soluble in 16 parts of boiling water, but deposited on cooling

Pyromeconic or pyrocomenio seid contains no oarboxyl-group, and its acid characters are very feebly marked. It orystallises in prizms, is readily soluble in water and alcohol, melts at 117°, and boils at 227°, but sublimes slowly at the ordinary temperature and readily at 100°

its positive detection is a decided proof of the presence of a preparation of opinm. It is not poisonous

The microscopic appearance of the precipitates produced in not too didute solutions of mecome and or solutile meconates by barrum chloride, calcium chloride, potassium ferrocyanide, and hydrochloric acid are highly characteristic

The most characteristic reaction of meconic acid is the formation of a deep purplish red coloration on adding ferric chloride to the solution of mecanic and or a mecanite. The shade of colour is distinctly different from that of the feiric acetate or formate. and the ferric meconate also differs from these in not being readily destroyed by boiling, or by adding cold dilute hydrochloric acid. and from the ferric thiocuanate in being unaffected on addition of mercuric chloride or autic chloride 1 If any doubt exist as to the presence of an acctate, it is desnable to precipitate the neutralised solution with nitiate or neutral acctate of lead, wash the precipitated lead racconate thoroughly, suspend it in water, and decompose it with subhuicited hydrogen. After evanorating the filtered hound at a gentle heat to drive off the excess of subhuretted hydrogen, the test with ferme chloude may be safely applied Instead of adding ferric chloride to the solution of meconic acid. the reagent may be applied to the solid substance, as obtained by the evaporation of its aqueous or ethereal solution

The red coloration produced by meconic acid and a ferric salt is much weakened by oxahe and phosphoric acids, and still more so by metaphosphoric acid

Comenic and pyromeconic acids also stilke a red coloration with ferric chloride, but with the latter acid the colour is less deep

Meconic acid may be extracted from its acidulated solutions by agitation with after, a property which enables it to be readily separated from morphine, acetic acid, tainin, and other substances hable to interfere with the observance of its reaction with farmos chloride. The extraction is not perfect, even when several more repeated, and hence the method cannot be employed for quantitative purposes.

Mocome and may be determined by converting it into a lead sall, or colorimetically by ferric chlorids, by comparing the depth of turn produced by the sample with that obtained by treatment with a known quantity of opium. Very fair approximate estimations of mecome and, and less accurately of opium, may be made in this way, even when the quantity of material at disposal is very insignificant.

Three of the atoms of hydrogen in meconic acid are replaceable

¹ Theoryanates (sulphoryandes) exist in sensible quantity in the saliva (and hence in the contents of the stomach) and also in white mustard. by metals, but recent researches have shown that the and is, properly speaking, dibasic, only two carboxyl groups, CO OH, being present. The third atom of hydrogen belongs to hydroxyl, and when this is replaced by metals basic saits of a yellow colour iesuit

The metallic meconates are mostly insoluble in water, except the meconates of the alkah-metals. They are nearly all medulic in alcohol, and are but slightly affected by acetee acid. The saits having two atoms of bease hydrogen replaced by metals are neutral to likuus anear.

Acid Calcium Meconate, Call_C, H(OII)O₁, as preceptated as a sparnelly soluble salt of characteristic nucressogne appearance on adding calcium chlorule to not too dilute a solution of meconic acid or a soluble meconate in presence of free ammonia, the solution of the solution, yellow, decake meconate, $C_{\rm sol}(Z_{\rm sol}(DII)O_{\rm sol})$, as precipitated On treating either of these salts with hot dilute hydrochlame acid, meconic and enjatallies out on cooling

For Meconates Ferrous meconate is a colourless, very soluble salt, which turns ied on exposure to an. Ferric meconate exists in the purple-ied liquid produced on adding a fairic salt to a soluble meconate

Lead Moonate is obtained by precipitating mecomic and or a meconic (or an aqueous solution of opium) with neutral acctate of lead. The triplimible meconate is stated to be formed even in presence of axcess of meconic and, but it is more probably a mixture or compound of the normal meconics, PfC-LLO, with lead hydroxide. The precipitate is quite insoluble in cold and boiling water, and is not affected by acctace and

Morphine Meconate has already been described (page 313)
Action of Solvents on Optim

The action of different solvents and reagents on opium and its constituents is shortly as follows —

Water dissolves mesoune and readily, as also subhate, meconate, and acetate of morphine The morphine is very spanningly soluble in cold water, and narcotine still less so Nascome is much more soluble than morphine, while the resm, caouthouc, &c., are mostluble, though certain gummy matters sees into solutions.

Alcohol dissolves free morphine as well as the acetate and meconate The other alkaloids of opium, as also the resun and caoutchouc, are dissolved by alcohol

Amylic alcohol dissolves all the alkaloids of opium, if in a free state The resun also is slightly soluble in amylic alcohol

Ether, benene, and earbon disulphide dissolve only about 05 per cent of free morphine, but the other free alkaloids of opium more readily. These solvents also dissolve the caoutchouc, but not the resin Acids dissolve all the alkaloids from opium, together with a resinoid substance

Fixed alkalies, used in excess, dissolve morphine freely, while narotine remains insoluble. Lime water dissolves morphine, but is a solvent for naicotine only in presence of morphine. The resin of optium is partly soluble in alkalies.

Ammonua dissolves morphine sparingly, narceine and codeine readily, while the other alkaloids and the resin of opium are moduble

From the foregoing statements, the arrangement of which is mostly due to E L. Cleavei (*Year-Book Pharm*, 1876, page 502), it follows that an *aqueous solution* of opurm will contain sulphate and meconate of morphine and other alkaloids, calcium salts, meconic acid, extractives, and resinous matter

An alcoholic solution will contain, in addition to the above, free narcotine, caoutchouc, fat, and resin

Oppum which has been exhausted with water still retains a bitter taste, but this is probably due to narcotine, as it is removed by carbon disulphide, benzene, or ether, in which morphia and its salts are insoluble. Water, even when cold, may be trusted to dissolve the whole of the morphine from opium, if the resultant solution be distinctly acid. In some processes of assaying opium, the sample is subjected to a preliminary treatment with benzene, chlorotorm or ether to remove narcotine, caoutchouc, and colouring matter (see page 349) By this means the subsequent exhaustion with water is much facilitated, and a puner solution of morphine is obtained In presence of much narcotine, morphine is soluble in benzene, but this is not true of the sulphate, meconate, or other salts of morphine Hence there is no loss of morphine on extracting opium with benzene. Meconate of morphine is, however, freely soluble in a mixture of alcohol and chloroform, but the simultaneous presence of ether prevents its solution more or less completely

ADULTERATIONS AND ASSAY OF OPIUM

Optum is hable to a varnety of adulterations, some of which are of a very gross kind Sand, chy, ashes, stones, shot, builts, lead tunnings and other maker-weights are occasionally mel with Singar, gum tragscanth, bulp of apmosts and figs, pounded poppy-capsules, and other vegetable substances of a socclaime, muchingmous, and resimous nature are also employed. Aqueous extracts of poppies and of Glametine latean are said to be sometimes added in Turkey, though rarely if ever seen in the opium imported into England. Such adulterants are industed by the darker colour and hygroscopic character of the product, by the difficulty in filtering the solution, and by the continuous strake which the simple leaves when drawn and by the continuous strake which the simple leaves when drawn

across a sheet of paper, whereas good oppum makes an interrupted mark

The proportion of safe yielded by opium should not exceed 8 per cent. The proportion of water in opium averages about 20 per cent, the usual range being from 15 to 28 per cent. It is best determined by taking a known weight of the opium in thin slices, and noting the weight on drying at 100°.

The extract of opium is determined by exhausting the drad sample with cold water, and collecting, drying, and weighing the sendue, or evaporating the whole or an aliquot part of the soluion to dryines, and weighing the extractive matter left. Should the insoluble resultue exceed 40 to 46 per cent of the drad sample (equal to a minimum of 55 per cent of extract), the presence of sand, clay, or other insoluble (mineral) adulternite is probable, while if the residue is below this proportion the presence of sugar, gum, or other soluble impurity is indicated?

According to Hanbury and Flinckiger, desed opuum from Asia Minor should yield from 55 to 66 pa cent—generally more than 60—of extractive matter soluble in cold water, the proportion of extract from Indian opuum boung from 60 to 68 per cent

The following are determinations by D. B. Dett [Tast Rock Phases, 1876, page 488) of the leading constituents of eighteen samples of optum, purchased from designate of good standing in London, Doblin, and Edinburgh. The squeenes extract was determined by subtracting the sum of the water and mobilial matter from 100 00. The proportion of morphia calculated on the dased optum averaged 11 06 per cent. The proportion of morphia in the div extract was 188 are cent (compare mage 389).

					1	ercentage Compo	Percentage of	
	Description	of Op	um		Water	Insol Residue	Aqueoue Extract	Morphine (hydrated)
1 2 8 4 5 6 7 8 9 10 11 12 18 14 15 16 17 18	Turkey,	· .	:		19 6 20 0 20 0 21 2 22 0 18 4 27 2 21 2 21 2 22 2 21 4 14 0 15 2 18 6 14 8	22 00 28 85 25 95 23 70 30 96 25 46 25 46 25 46 25 80 38 80 29 70 47 90 26 80 27 40 26 10 27 40 26 10 27 20 28 20 29 20 20 20 21 20 22 20 23 20 24 20 25 20 26 20 27 20 27 20 28 20 20 20 20 20 20 20 20 20 20 20 20 20 2	47 80 51 15 48 06 55 10 47 06 50 15 54 90 45 40 47 50 20 90 50 20 00 60 00 70 01 20 56 90	10 75 112 90 10 20 7 67 0 00 11 99 12 90 6 76 6 76 9 90 8 85 0 093 6 100 8 60 2 100 7 7 7 80 5 88 7 7 90 6 7 80
10.		rage,	•		19 70		50 44	8 88

Hager recommends the following additional tests for the punty of opum -25 grains weight of the previously dried sample is tritmated with half an ounce of boiling water, when the formation of a stiff maste will indicate the presence of starch, flour, com. &c. 2 ounces of water should next be added and the hound filtered If the filtrate be brown or of a deeper colour than "wine-vellow," the presence of liquorice or pther vegetable extracts is probable. The liquid should have an acid reaction, or admixture with chalk, lithage, or ashes may be suspected. The hand should give no reaction with potassium ferrocyanide (heavy metals) and if evanorated to one orner and treated with twice its measure of alcohol no precipitate should be produced (indicative of adulteration with gum or ceitain salts)

On agreeting powdered oppun with chloroform, any starch or mineral adulterants will settle out, and may be weighed and further examined microscopically and chemically

When moist, opium is very liable to become mouldy, and hence should be died at a moderate temperature and carefully preserved from the air. If kept in a damp condition, fungoid growths soon make their appearance, and gradually diminish and destroy the

groma of the onium, besides materially reducing its alkaloidal value DETERMINATION OF MORPHINE IN OPIUM, MORPHIOMETEY,

By far the most important item in the examination of onturn is the determination of the morphine present. The proportion of this constituent values considerably, as already stated; but dried and powdered oppum intended for medicinal use should not assay Less than 10 per cent 1 This is the limit of the German and Austrian Pharmicoposias, while that of the United States allows the range of 12 to 16 per cent, any nicher oppum to be reduced within these limits by mixing it with an article of lower grade in proper proportion. According to the German and United States Pharmaconomias, ontum in its normal moist condition should yield not less than 9 per cent of morphine The British Pharmacopona of 1867 allowed a range of 6 to 8 per cent., but in the edition of 1885 the assay is directed to be made on the dired substance. the yield of morphing to be between 94 and 104 per cent 2

The difficulty caused by the natural variations in the quality of opinin is well met by a process patented by B S Proctor, who removes the greater part of the fatty and resmons matters and the worthless narcotine, and reduces the opium to a uniform rectified condition, in which it contains 10 per cent, of moinfune

^{2 &}quot;This standard is ridiculously low, and will have the effect of depriving medicine of all the best opium that reaches this country. This standard is about equal to that of the last Pharmacopena, but then there was no man-

The assay of opum for morphine has received much attention, the investigators bung very numerous and the biblography very extensive. The accurate determination of morphine in opinus is attended with peculiar difficulties, and many of the processes which have been published give little better than rough approximations to the truth, especially when employed for the assay of abnormal samples! Of the many methods proposed, the following are among the best —

British Pharmacopensa Process 2—This method of assay is basted on —the conversion of the resunous matters of opuum tuto unabled hime compounds, the decomposition of the morphise measurable with formation of menabolise calcium mercentse, the solubility of the resultant free morphise in line-water, the decomposition of the solution by ammonium chloride, with formation of calcium holoride, ammonia, and free morphise, the use of alcohol to dissolve impurities, and of either to promote the crystallisation of the akiladid, and the collection, weshing, and weighing of the morphise thus obtained The following are the details of the process as land down in the Buttish Pharmacongus of 1885.

"Take of powdered opum, drued at 212" F (=100" C), 14c or grams, lime, freshly slaked, 60 grams, shorded of ammoutum, 40 grams, rectified spirit, ether, distilled water, of each a sufficiency. Triturate together the opum, lime, and 400 grams—neasures of distilled water in a mortar until a uniform mixture results, then add 1000 gram—neasures of distilled water, and sirr coessories, and 1000 gram—neasures of distilled water, and sirr coessories, and all 1000 gram—neasures of distilled water, and sirr coessories about 5 miches in diameter into a wide-mouthed bottle or stoopered about 5 miches in diameter into a wide-mouthed bottle or stoopered.

name standard green. It is all very well to standard me repranhons, but, I think; it is going too far when we attempt it with actual products, but if we no to have a maximum and minimum standard for opium, let it be one which will include the best and crolleds the inferen and salidarated kinds, instead of the reverse, as one obtains. To statam this it would be nocessary to save the standard of least 2 pn cent "—(Michael Conroy, Phons Jose, [3], xxx 378).

¹ The sampling of opium for the purpose of analysis is not always an easy operation, and is not conducted on a uniform plant. J B Nagel worst recommends that a small since should be out by a lamfs from the interior on each imp of the lot, these poses mixed together, and 10 gammes taken for the determination of mostime. The remainde is dised, pulvenised, and the cushind most in each more him for the property of the determination.

² This method was originally dovased by Portos and Langlors (Okra-News xir 67), and with slight alterations was adopted by the Société de Pharmace of Parrs, and made official in the United States Pharmacopcial of 1850. It was further unproved by M. Contoy (Pharm. Joss., 13), xv. 474), and adopted as the official pear in the Britals Pharmacopcian of 1855.

flask (having the capacity of about six fluid ounces, and marked at exactly 1040 grain-measures) until the filtrate reaches this mark 1 To the filtered liquid (representing 100 grains of opium) add 110 grain-measures of rectified spirit, and 500 giain-measures of ether, and shake the mixture, then add the chloride of ammonium, shake well and frequently during half an hour, and set it aside for twelve hours.2 Counterbalance two small filters, place one within the other in a small funnel, and decant the ethereal layer as completely as practicable upon the inner filter Add 200 grain-measures of ether to the contents of the bottle and rotate it, again decant the ethereal layer upon the filter, and afterwards wash the latter with 100 grain-measures of ether added slowly and in portions. Now, let the filter dry in the air, and pour upon it the liquid in the bottle in portions, in such a way as to transfer the greater portion of the crystals to the filter. When the fluid has passed through the filter, wash the bottle and transfer the remaining crystals to the filter, with several small portions of distilled water, using not much more than 200 grain-measures in all, and distributing the portions evenly upon the filter. Allow the filter to drain, and dry it, first by pressing between sheets of bibulous paper, and afterwards at a temperature between 131° and 140° F (55° and 60° C), and finally at 194° to 212° F (90° to 100° C). Weigh the crystals in the inner filter, counterbalancing by the outer filter crystals should weigh 10 giains, or not less than 91, and not more than 104 grams, corresponding to about 10 per cent of morphine in the div, nowdered onium"

The skilled chemist will find alundant opportunity for improving on the method of manipulation presented in the above process. It will probably substitute their equivalents in grammes and estimaters for the weighted and measured grams prescribed, but he will, in practice, find at advantageous to increase the weights of opium and lime taken to 10 grammes and 5 grammes respectively, and the measure of the water to 100 cc 52 cc of the filtrate will then represent 5 grammes of the opium, and the delay, consequent on collecting so large a poiston as \$6\$ of the

¹ The additional 40 grain-measures is intended as an allowance for the average increase in the volume of the liquid caused by the extractive matter of the column.

² "The use of an excess of ethes, much keyond either saturation, so as to cause an ethecial layer to two shorts the crystallising hinds, along with the flequent shrking mp of the ether with the aponent liquid; an interest clarking crystallisations, marks an important advance in opium saxy" "(-6.8 B Pres not t;) The mactice has been adopted in all recent methods of assaying opium.

liquid, will be avoided. A less clumsy means will be adopted for measuring the exact quantity of the filtrate required than that of relying on a mark made on the side of a 6 or bottle, on the broad part of a flask, and the othereal layer will be removed by some form of pipette unstead of attempting to decent it on the filter.

The BP, process for the assay of opum is tolerably simple and rand, and when cascidity executed, gives fanly constant results. As suggested by Conroy, and proved by Banthwate and Farr, the time allowed for precipitation of the morphine may be reduced from twelve hours to two without affecting the accuracy of the results, but it is safet to allow six or eight hours to elapse before filtering. It would be a further miprovement to direct the behavior of the provided of the provided and the properties of the would be a guarantee of the true nature and purity of the precipitate, and would save the time required for, and uncertainty attaching to the drying of the alkaloid

The results yielded by the B.P process of assaying opium are seriously below the truth, a fact ignored by the editors, although pointed out by M. Conroy, whose process it practically is ⁸

Braithwaite and Farr (Pharm Jour, [3], xvn. 398) confirm Conroy's view, and state that the morphine left in solution is about 1 per cent of the opium But they point out that the precipitate contains an average of 7 per cent of colouring matter

² M Conroy states that, by reducing the quantities of opium and water recommended by him, the editors of the Phirmacopum have deprived the process of one of its chief ments, with the consequence that the 1040 grammassizes of filtrate required can only be obtained at the secrible of much time. A O A brahem (Pharm. Jour. [2], xm 280) endouses this view holding that: "for the sake of saving a few grains of jouinum, a numple and quick process had been inedered most technous. The standard of 10 per cent was, processes, and the same to the control of the same of the same of the standard of 10 per cent was, processes, and the same to the same of th

²Titration of the prespitated morphine was directed by Portes and Langlois, the original propossis of the method (Jour Pharm et Chemie, November 1881)

According to the Pharmacopous, from \$4\$ to 104 grams of crystals should be actually obtained, "corresponding to about ten pae ent of morphans in the dired produced or pum," a statement which is materially uncourse to our roy found, in test-expansances on 10 grams of pure morphins, 906, 903, and 908 grams were scovered, thus showing a notable but almost constant loss. The loss when an aqueous orchard or journs in operated on, unstead of a pure solution of morphins, is still greater, probably ranging from 1 to 12 per cent. Hence a yield of \$9\$ to 10 per cent of morphins, by the B P process, not unprobably corresponds to about 11½ per cent of morphins charge from the morphins actually present.

as imparity, and hence, in assaying an opium containing 14 to 15 per cent of maphine, the eron from this cause approximately balances that due to imperfect precupitation. On dissolving the impure morphine in hime-water, a large proportion of the colouring matter is left in this filter, and on extracting the solution with ammanium chloride, alcohol and either, as in the B P process, the rest of the colouring matter remains in solution, and the representation opinion is obtained almost white. But there is a serious loss of 10 per cent of the weight) through solublivity of the precupitate

J De n la m Sm 1 h (Chem News, 17m, 93, 163) obtained by the B.P. process, in five experiments, results ranging from 94 to 90 per cent, a sixth experiment giving 10 b per cent, the true amount of morphine present being stated at 112 per cent, which was obtained by a process giving exceptionally high results (page 347) Smith distauts the use of lime as open to many objections, and this opinion is sheared by R Willia ms 2 (Chem News, 17m, 134), who gives the following results obtained from four samples of opinion when assigned by the processes of the British, Amenoca, and German Pharmacopeans respectively

	No 1	No 2	No 8	No 4
Britseh,	108	10 5	7.4	12.2
Amesican, .	11 1	108	81	119
German,	10'2	10-0	71	106

In each case the German process gave the lowest result, and the American the highest, except in the case of No 4 sample

Dott considers 7 per cent of impurity excessive, and thinks 3 to 5 per cent would be nearer the first.

"Notwithshudding this, D B Dot's (Pharm Jone, 13), xxx 33) considered that the endpoyment of lum "in an unch to be and un the force in Term a power solution of morphine than can be obtained by any other single operation, and benefic climinates nearly 21d possible additionants. The increase in the properties of the summonium chloride is usually semi-habity jume, we may always if the option in genome Sample sue, however, occasionally mark with which yield with the chloride of ammonium a cestion amount of forceding receptible along with the morphism. In such cases it is presummently necessary to apply the thirston with standard and Those can be no doubt that the althos of the Pharmacopiers ought to have allowed for the meritable loss of morphism on the mather waters, supexually when any other trustworthy methods in particular.

⁸ Various observers agree that the results obtained by the German method of sasy are at least 2 per cent, below the tutth, and the morphine not always pure (Pharm Jour., 131 xir 645).

The yield of maphine obtained by the B.P process ought to be corrected by a definite allowance, but a more satisfactory has would be to prescribe a method by which the remaining alkaloud could be recovered if desired. This might probably be approximately effected by agriating the warm ammoniscal filtratic with anythe alcohol, and separating and evaporating the solvent

United States Pharmacopens Process—As already stated, the method of opum assay presented by the Dratia Pharmacoperia as a modification of that previously adopted in America The latter differs from the BP process chiefly my prescribing the use of a larger proportion of ammonium chlorule. This is a distinct disadvantage as tending to retain morphine in solution, a fact pointed out by M Conroy, and confirmed by Wrampelmorer and Meinner!

Method of Teachemacher and Denham Smith—These chemists have examined most of the published methods of assaying opium (Othem News, Ivi 193, 103), and have found them waiting in one or more respects. They reject methods in which the precipitation of the morphine is effected in presence of more than a very limited amount of alcohol (a.g., Flinkhyer's dolle methods) as likely to yield low results, though a very pure product, they object to the use of hime (as in the BP product) as causing the product to be coloured, and being open to other objections, and they strongly advocate the titiation of the morphium isolated, maked of determining it gravimetrically. All these objections are well founded, though searcely so ritle at they are regarded by the authors, who, however, have described a method of assay which, on the whole, is probably the best inthacts published? The process they recommend is

¹ These latter chemuts calculated the amount of ammonum chlorde whole would remain in excess, and free ammons which would be produced in the reaction, and successful them solvent action on morphine, but the correction logically based on their results would be sensowly in excess of the sciental love of morphine in pactone. H. Li by d has proposed to correct the results of morphine in pactone. H. Li by d has proposed to correct the results of morphine in pactone. H. Li by d has proposed to correct the results of the most price of the process of the process of the sensor of 1 per cent to the morphine thus found Than, # 0 per cent of morphine he actually recovered, according to H. Libryl, the trans amount present is 9 of 2, 10 e+1, 0 community and according to H. Libryl, the trans amount present is 9 of 2, 10 e+1, 0 community and according to the part of the part of the process when the process of the part of the par

² This view is confirmed by D B Dott, who operates as follows:—10 gammes weight of the powdered opinm is exhausted with proof spirit, one or two drops of ammonium oxilate are added, and then ammonia, until the spirit so only slightly said. The legand is then evaporated to one-third, allowed to

founded on one originally devised by Prollius and modified by F. A Fluckiger (Archiv des Phasm, [3], xxvi) It was then materially improved by E R. Squibb (Ephemeris, 1 14), and assan further modified by C. M Stillwell (Chem News, ly. 41. 54) The following are the details of the process as prescribed by Teschemacher and Smith -200 grains weight of opium is thoroughly exhausted with waim distilled water. and the hound filtered. The aqueous extract is concentrated to a thin syrup in a shallow dish, over a water-bath, which by proference should not boil. The syrup is transferred to a suitable flask, and the dish washed out with a few drops of water. To the contents of the flask are added 50 fluid grains of alcohol (specific gravity \$20) and about 600 fluid grains of ether A soft cork is inscribed and the contents of the flask mixed gently but thoroughly, after which 50 fluid grains of ammonia (specific gravity 935) should be added The flask is then well shaken to precipitate the alkaloid in arenaceous crystals, and occasionally agitated during the ensuing eighteen hours. The contents of the flask are then transferred to a vacuumfilter, and when all the adherent liquid is diawn out the crystalline precipitate is washed with "morphiated spirit" until the liquid passes through colourless It is then washed with "morphiated water "2 until this also passes colourless. The precipitate is then dried, at first slowly and afterwards at 100° C The dried substance is then finely powdered and digested thoroughly in benzene to dissolve the narcotine and such other opium alkaloids as may be present in addition to morphine 3 The liquid is filtered and the cool, and filtered. The filtrate is concentrated to about 5 co , transferred to a small flask, and the cansule washed with 4 c c of water and 3 of methylated spirit Next add 2 2 o.e of ammonia solution (sp gr. 960) and 25 cc of other, and agreate After 18 hours, decant the other as completely as possible, receive the aqueous liquid on a counterpoised filter, wash with morphiated water, dry, wash with benzene, dry, weigh, and tatrate the whole or a portion with decinormal sulphune acid (Pharm Jour , [3], 1xii 746)

Rowland Williams digests with cold water for twelve to fourteen hours, and claims to obtain a cleaner solution than when warm water is used.

² The "Mosphatot Sphrit" is made by mixing 1 measure of ammonia (epecing garrier 880) with 20 of methylated sparin, and digesting in the liquid a large excess of powdered morphane for several days, with frequent agrisance. The little of larged contains 0.3 per cens. of morphina, "Mosphated Wades" is made by agitating cold water with excess of morphina, and filtering after twenty-found hours. The filtitudes contains 0.0 to go cont. of all/acide.

Sexing that the morphino is ultimately determined by titration, that marcoine, ancience, and papaverine have no action on himms, and that codence is solitule in 80 parts of cold water and readily solitule in alcohol and either, the presented treatment with benzaon in order to remove these alkaloids esents superfluors. When the morphine is to be weighed, it would probably an event superfluors.

precipitate further thoroughly washed with beazene. The reading will consist for incryline "free from other orquin alkaloids and nacotine, but still containing coloring and possibly other corgame matters to the extent of 3 to 10 per cent." (of its weight) The powder is dired, weighed, and thraded with limins and a standard hydrochlorine and, peparate so that 1000 gunus by weight will exactly neutralise 100 grains of pure morphise crystal-lised from water, washed with eithe, and gunty dired at 100° C

Fluckings's process F A Fluckinger has devoted much attention to the assay of opium, his most recent method (Archiv Pharm, [3], xxvii, 721, 769, Pharm Jour, [3], xx 588) being as follows -8 grammes weight of powdered opium is placed in a plaited filter, and dried at 100° for half an hour. It is then treated with 20 e.c. of a mixture of equal measures of ether and chlorofoim, and when this has run through, with 10 cc of unmixed chloroform. The filter and its contents are then dried at a gentle heat, and the powder vigorously and repeatedly shaken in a flask with 80 cc of water to which 0.2 gramme of ammonium oxalate has been added After two hours the liquid is passed through a dry filter, and 42 5 grammes of the filtrate (=4 grammes of sample) treated in a small tared flask with 71 cc of rectified spirit, 15 cc. of ether, and 1 c e of ammonia (specific gravity 0 96) The mixture is frequently shaken during six hours, after which the hound is noured on a double filter, the flask runsed with 10 cc of water or morphiated water, and the rinsing used to wash the filter precapitate and inner filter are dried at 100°, returned to the dried flask, and the whole further heated to 100° tall constant, the outer filter being used as a counterpoise. The foregoing process would be materially improved and shortened by titrating the dried morphine instead of weighing it on a counterpoised filter, and its accuracy increased by reducing the quantities of liquid used J B Nagelvoort (Pharm. Jour, [3], xx1 598) has slightly modified the above method, which he commends very highly, for the assay alike of opium and its galenical preparations. He found the isolated alkaloid to be completely soluble in 100 parts of lime-water to a clear, colourless solution, whereas the "morphine" obtained by Squibb's and Stillwell's modifications of Flückiger's former process contained from 10 to 20 per cent of impurities 1

be better to wash with morphiated spirit only, and when it is to be titrated to omit this treatment and wash it at once with benione

If natorine be present it is left as a crystalline residue on testing the sikaloid with hime-water. Perget has proposed to purify morphine by dissolving it with dultie acetic acid and adding potavium ferrogrande, filtering, and precipitating the mosphine from the filtrate by ammonia.

L. Kı offer, ın 1887, described a volumehre process of easying opuum, besed on the reaction of the monphine with potassum forregrande, reaction of the excess of this sait with potassum include, and tetestion of the blerated uotine with standard thoushplate (Annal Chem Pharm, un. 189). A limited number of experiments made in the similar's alboratory on Kieffer's process have not valided encourancing results

Extract or Orius, \hat{B} P, is made by exhausting the 'opinim' with sold water, straining, and evaporating the liquid to half the weight of the opinim used. It has a public consistency, and is said to yield about 20 per cent of mergline when assayed by the official test for opinim. W. P. W. ant. (Pharma Jony; [3], xvi 989) found by this process from 9 to 804 per cent of morphine in ax samples of the commercial extinct of opinim? By the method of the 1867 Pharmaconous, D. B. Dett found in eleven samples of extinct, purchased from drugasts of good standing in London, Dublim and Eduburgh, proportions of another analysis of the 1867 per cent, the mean being 100 per cent.

Laqual Extract of Opnum, BP., as prepared by macerating I ounce of the solul extract with 16 ounces of water, adding 4 find ounces of rectified spirit, and filtering. It should contain "22 grains of the solid extract in nearly 1 fitud ounce." The specific gravity should be between 0 985 and 0 995, and when as-ayed by the process prescribed for opnum "should yield about 1 per cent of morehine"

J. We od llanu I (1"as-Book Pharm, 1882, p 514) found in ten sumples of the hund extract of quium of commore propunctions of solid readule ranging from 3 02 to 4 92 per cent, and of morphane from 0 19 to 0 37 per cent Tisses detasimations were made by a modification of Piollius' mothod, the securacy of which was demonstrated. D B D ot 1 (Yas-Book Pharm, 1876, 500) found the specific gravity of elevent samples of commercial fluid extract to range from 0 985 to 1000, while the proportion of morphine per fluid ounce varied from 1 66 to 451 crains

TINOTURE OF OPIUM LAUDANUM For the preparation of this important medicine, the Bistish Pharmacopous directs to "macerate 1½ ounces of opium in powder in 1 pint of proof spirit for

¹ The United States Pharmacopous orders an addition of 5 per cent of

² J. H. Hoseason (*Phus m. Jour.*, [3], xix. 764) has pointed out that extract of opium is sold by wholesals druggists at a cheaper rate than they can purchase the opium for its preparation

seven days in a closed vessel with occasional agriation, then strain, press, filler, and add sufficient proof spirit to make one pint I foundants the soluble matter of 33 grains of the opium, nearly, in I fituid ounce, or about 33 grains of morphine in 1 fituid ounce, and about 33 grains of morphine in 1 fituid ounce, and the solution of the property of the solution of the property of the solution of the property of the solution of the strain of the property of the same of the solution of the size of opium and the property of the same of the size of the size of the same of the size of th

W P Want (Phorm Jon., [3], xr. 959) found the specific gravity of vs. samples of incube at optum pocured from leading wholesalo houses to tange from 931 to 939. The proportions of morphine were estimated (in disphase) by the official process for optum (using about 3 ounces of the tracture), and were found to be respectively — 33, 33, 26, 33, 34, and 218 grams per fluid onne. All six samples were very similar in appearance and odors.

J H Hoscason (Pharm Jour, [3], xix 754) has published the following results of the examination of ordinary commercial samples of fracture of opium —

¹ This statement of the condition of evistence of the mosphine is without warrant, and is opposed to the known facts. The very existence of "binno-conate of morphine" is doubtful, and a large propertien of the mosphine in optim exists as sulphate.

⁵ The Tincture Open of the United States Pharmacopera (1889) is prepared from powdeed opms (asseying 12 to 16 per cent of morphise) 10 patts, water 40 parts, alcohol (specific gravity '820) 18 parts, and sufficient dilute alcohol (specific gravity '925) to make the twicture obtained by percolation up to 100 parts. All the ingredients are by weight.

The Tuntus Oyu. Sumples of the Germ an P harm as op nata (1869) is prejucted from powdered opinum (with 10 per cont or mass of most phune) I part, diluted alcohol (spendic gravity 80 to 856 at 18' 0) 5 parts by weight, and water 5 parts. It has a spendic gravity of 79' at to 97's, and contains, in 100 grammes, the solible portion of heavily 10 grammes of opinum, or approximately 1 per court of morphise. O grammes when assayed should yield not less than 0.38 gramme of mosphise.

The corresponding preparation (Tractura Extractor Opis) of the French Codex (1884) to prepared from 10 parts of extract of opinin (-16 7 of dry opinin) orbitaining 10 to 12 per cent. of mosphine, and 120 parts by weight of alcohol of 912 specific gravity

From these particulars it is evident that the strength of the official tinctures of opium vary convolciably, both in alcoholic strength and the proportion of morphine. The United States and French preparations are the strongest (in alkaloid), the German weaker, and the British the most duties.

Number	Specific Gravity,	Absolute Alcohol, per cent by weight	Residue; grs per fluid ounce	Motphine, grs ps: fluid onnce	
1 8 6 6 7 8 9	964 962 912 940 962 960 961 961 960 961	28 40 41 42 30 87 87 86 88	18 6 17 0 20 5 14 5 15 3 17 3 16 0 13 5 14 6	8 % 8 0 6 0 2 0 2 0 2 0 2 0 2 5 2 7 8 0	
Avelage,	955	88 4	164	9.8	

Six of the above samples were evidently made with a mixture of equal measures of rectified spirit and water, instead of the proportion of 5 3, which would yield approximately proof-spirit.

J Woodland (Ysen-Book Phanm), 1882, mag 514) found the solid results from fourteen samples of ancture of optum procured from both London and provincial chemists to range from 321 to 511 per cent; while the morphine (estimated by a modification of Prollins' method) ranged from 0.3 to 0.70 per cent.

D. B Dott (Tax-Book -Planm, 1876, page 560) found the specific gravity of twelve samples of the communical nucleus of epum to range from 922 to 962, while the crude morphine (estimated by a modification of the BP 1867 method, are regularly τ_{ij} of pure alkadoul) contained in the same spectiment and ax others (the density of which was not observed), need from 4.37 to 0.55 grains per fluid ounce, the average being 2.96

From the foregoing published results it is evident that the composition of commercial functure of opium varies to a very discreditable extent, both in alcohole strength and the proportion of morphime contained in it. Still greater variations in strength are to be found in the structure when purchased under the of "landamum," which, however, is now an official synonym for function of committee of committee

S. J. Hinsdale (Chem News, lxu. 77) has described a simple

¹ Several presentations have occurred under the Sale of Food and Diago, Act for the sole of delective incitate of opium. In the case of White a Phywatzi, it was sold under the official mane to the written order of a metical man five outer accepted the view of the defence, that, as the preparation contained alsohol, it was a "function," and that if it contained any opium at all it was a "function of opium," which, consequently, might be of any strength whatever This deceason was reviewed on appeal to the Court of Queen's Bench (Pharm Jon. [13], 3rt 1960)

method of determining the morphine in tineture of opium by observing the depth of the blue or green coloration produced on treating the sample with a freshly prepared muxture of ferric chloride and potassium ferricyanide solutions

COMPOND TINGTURE OF ČAMPHON, BP, as the formal designation of the prepandion popularly known as "Parsgolie" or "Pargolie" Elixii". These names were adopted as official eynonyms for compound instairs of camphor in the reprint of the British Planmacopoins of 1886, and hence preparations sold under these titles ought now to be strictly of the quality and strength of the BP undrum Compound instairs of camphor is directed to be prepared with 40 guins each of opum and beneon each, 30 grains of camphor, and 30 minus of oil of anne, the whole being diluted with proof-spirit to 20 fluid numers.

Much of the paragone or compound tracture of camphor of commerce is deficient in one or more of the constituents. The spirit being the most costly ingiculent, there is a strong inducement to the vendor to reduce its amount, a practice which is objectionable because the prescribed proportion of oil of anise cannot be kept in solution in a very weak spirit. Sometimes only traces of oil of anise are present, in which case the finetine remains clear when diluted with three or four measures of water The benzoic acid is sometimes deficient in quantity, and occasionally wholly absent, even in the case of tinctures purchased from registered pharmacists. The opium is the most important constituent of paregoric elixir, and is apt to be deficient in amount or quality, besides being frequently wholly omitted. The last practice is due to the fact that preparations of opium cannot be legally sold except by registered pharmacists, and hence a preparation destitute of opium is largely substituted by general shopkeepers for the genuine "paregorie" or "compound tinctuie of camphor" sold by the dauggests 2 In an anstance within the personal experience of the author, the opium of paregoric clixir was replaced by henbane Potassium and ammonium bromides are extensively used in factitious paregone

The proportion of alcohol in compound tineture of camphor is

¹W D Mason (Pharm. Jour, [3], xm. 396) points out that great saving of time and torolle in mecention, agritation, siltering, &c, could be effected, and a pea feetly clear and bught tuncture, previously the same as that of the Pharmacopeus, obtained by adding the optium in the form of a ready-made timeture

² So-called "panegone" is vended by costermongers in the streets of London

indicated with approximate accuracy by the specific gravity, which should not be higher than 0 926.1

If a measured quantity (35 cc) of paregone be rendered distinctly alkaline with soda, and evaporated to about 10 cc., the alcohol and a portion of the camphot and oil of anise will be volatilised On then shaking the liquid with ether, the remaining camphor and oil of anise will be extracted. If the ether be separated, and the aqueous liquid acidulated with hydrochloric acid, benzoic acid will in some cases be precipitated, but whether it separates or remains in solution, it should be dissolved out by agitating the acidified liquid with other. On allowing the separated ethereal solution to evaporate spontaneously in a small beaker, the benzoic acid is obtained in a state at to weigh ,2 but a better and more rapid plan is to repeatedly agitate the ethereal liquid with water nutil the washings no longer redden hitmus, add a little more water and a few drops of phenolphthalcun solution, and titiate the liquid with m caustic alkali (preferably baryta-water), which should be added until the aqueous laver acquires a pink colour, not destroyed by agritation with the ether. Each 1 c.c of alkalı 10quired represents 0 0061 gramme of benzoic acid. If 25 cc of the tancture has been employed, the number of milligrammes of benzoic acid found, multiplied by 0 35, gives the grains of benzoic acid per pint of the tincture. The meconic acid extracted together with the benzoic acid is too small in quantity to affect the result, but its presence may be detected and the amount roughly determined by separating the ethereal layer after the titration is complete, and destroying the pink colour of the aqueous liquid by a drop of dilute hydrochloric acid On now adding a drop of ferric chloride solution, the deep purple-red coloration characteristic of meconic acid will be produced

The detection of meconic acid in the above minime of course proces the presence of opium in the function. When this misor-mation alone is sought, the paragonic may be diluted in a test-the with proof-squirt till it is of a light ytellow colour, and a drop or two of solution of fearnechloride them sided. If opium he present more or less deep red coloration will be produced, owing the formation of meconate of more. By comparing the depth of red colour with that given by a stanisal tincture, a rough indication of the proportion of opium present can be obtained, but the amount of meconic acid in opium is too visible to allow of much

Where a more exact determination is required, it may be made by the method described in Volume I, under the head of Tinetines

² The author has occasionally observed the benzuer and thus extracted to have a distinct unious odom.

stess being placed on the result obtained. It sometimes happens that paragonic so coloured with cochineal or contains a variety of tamin, in which case the coloration with force chloride becomes obscured. On cautiously adding hydrochlore and, drop by drop, the colour produced by tannate of iron is destroyed, while this disto the unconstant pensats till considerably more each has been added to the unconstant pensats till considerably more each has been added

The proportion of optum in paregone is too small to allow of the ordinary method of detainining morphine being conveniently used; but tair results, sufficiently accurate for most purposes, may be obtained by volumetine or coloimetine application of treatment with processing force and option of the continuous processing strength and only of the continuous processing strength and of the color o

Toxicology of Opium and Morphine.

In whatever form or manner it may be administered, opium is found to act as a typical and powerful narcotic, and in excessive doses is fatally poisonous ¹

In a letten to the olded, D. W. m. Mo or e, late Surgoon-General, Rombry, pouts out the exagement shatement muck experings the ill effects of optum enting and emoking. Ho writes —"No one denies that the accessare use of optum—whother emoked, eakan, or druml—produces unjurious consequences; but so does access in the use of sparts, of lossig goes, or even of funt. —I am quite sure that the use of oppun, speaking generally, in more advantageous than delections. Autt-opumnsts asset that all using the drug in any time por might be a surface of the state of the

" The use of omam, even in excess, is norther so deleterious to the consumer nor so dangerous to his neighbours as the use of surrits to excess The onum eater or smoker . . . attains to a placed repose, which is very different to the excitement caused by sunits . . Many maladies for which oppum to used in the East have been attributed to oppum. Numbers of neonle suffering from all kinds of maladies are to be found in Eastern orium houses. But the people thus affected fly to opium for a relief to suffering, and visitors finding diseased people in the opinm-houses have ignorantly attributed the maladres econ to the use of opium . . Opinm pievents elemacansis or waste of tisue, and thus contributes to endurance of fatigue, as evidenced by the long distances Kossids travel in India, their only support being a small pill of onum, a number of which they carry in a tin box. This is evidenced also by onum being given to camels, in combination with other substances. when these animals are called upon for extraordinary exertions. Onum also enables persons to live on smaller quantities of food than they could otherwise do-in this respect it resembles tea. Thousands wore kept alive during Indian faminee who would have succumbed from want of food had not opium been available. There is also no doubt that opium exerts a prophylactic The poisonous effects of opum are essentially due to the morphine contained in it, and the symptoms it produces differ but little from those consequent on the administration of pure morphine, except that there is a greater tendency to convations, and in the latter cases the effects are usually manifested more rapidly than in the former, generally commencing in from five to twenty minutes if the moscon has been taken in solution

After possuing by morphino or opium, dimness of sight and relaxation of the massles, with drownness and stupo, are amulty the flust symptoms observed. At first the patient may be sroused without such difficulty, but as time goes out this becomes impossible, the drownness passing tuto complete come, often accompanied by allow and extended as the student of the large majority of cases the pugits are strongly contracted in the earlier siages, but lates, and when a faste teammation is approaching, they are often dishated. They are usually meanable to light Occasionally, aspecially with excessive doses of opium, there is vomitting, or even puging. The pulse is at first weak, quick, and irresculate, but afterprarts slowe and full

Foscaning by morphime or opsum often closely simulates alcoholic dunkenness, and, in the absence of a smell of opsum in the breath or vomit, it is often very difficult to distinguish between them Come, due to urremia, apoplexy, or violence, may also be mistaken for postoning by outum or its .repeatstone.

The dose of morphino necessary to destroy life is extremely variable. Infants and young persons are peculiarly susceptible to opium and its preparations. Death has been caused to infants by

offect against malaneus fevers, which effect is recognised, not only in the East, but also in the against districts of this country. That it relieves chronic painful maladies does not require proof

"Poople in the East will have opium—for with them it takes the place of this standards or insoche—and they will have it in spite of any-thing east-opeumsts may advance to the contrary. In opium they have a change, eastly carried simulate or anxionic, according as they may two it, and nothing the anti-opeumsts may say will prevent the use of opium it, and nothing the anti-opeumsts may say will grevent the use of opium exter county pursons in more deleticent than sendengthe dusg, for it trutted free more with the digestive expectation. Taking opium in the form of opium water county pursons in his improves. Seeking opium is the hierts handfill manner of opium called chronidar or chemico, and, after much experience and investigation, I regard account good as harpines, unless radialpul in to evosse and the vest magnity of those sums quarties, proceed to excess—16 Fertland Paces, March 10, 18011"

A. Swaine Taylor mentions a case of opium poisoning in which one pupil was contracted and the other dilated

jith, jith, jith, and even a jith of a gram of opuum, as also by a few drops, and even a angle drop, of tenture of opuum. On the other hand, children have recovered after doses of 1 gram, 5 grams, and 7 j grams of opuum, and after two teaspoontols of laudanum Half a gram of morphum coatesta hea proved fixed to an adult, but as a rule, the usual minimum fatal dose for an adult may be stated as 1 gram of a salt of morphum, or 7 grams of opuum. Personal habit, as in the case of opuum-eaters, and idiosyncrasy will of counse largely modify the show conclusion.

The post-motem appearances of poisoning by morphine are by no means well-marked. The stomach and intestines usually appear healthy. If opium itself has been taken, its peculiar and characterishes closur may often be recognised when the stomach is flat opased.² Congestion of the lungs and biam as most commonly met with, but these appearances are not invariable, and when they exist, afford no definite evidence of opium poisoning. The blood is usually ver full.

Besides opium itself, morphine and its salts, and the various official preparations of opium (eg, the functure and extact), there are various notetrums containing opium, which have not unfrequently been the cause of death, especially in the case of mfants, for whom opiates may be regarded as generally dangerous and unsuitable ²

- ¹ The author has observed an unmastrkable smell of opium in the contents of the bladder sixty hours after death by taking laudauum
- ² Sprup of Poppies is professedly a sweetened decoction of English or white poppy heads. It is of very variable strength, and is said to be sometime substituted by a nivitine of tructure or influence of ordering with simple syrup.
- Winslow's Scotting Syrup sometimes produces symptoms of unitotic porsoning it is said to contain about 1 grain of morphine and other option alkaloids in an ounce (Phan Jour, 18], in 975)
- Godfy ny's Omdual is stated to be a mixture of treacle and esserfins with 1 drachm of timeture of opium in 6 ounces. Half a tempoonful is said to have caused the death of an infant, and in the years 1888-67, fifty art deaths were recorded from its use, probably by its administration in excessive doses by ignormatine names.
- Hawkswooth's Mixtus a contains magnesium carbonate, rhubarb, compound spirits of ammona, sweet spirit of nite, oil of cassia, simple syrup, water, and other ingredients, with 1 part of tinctine of opinion in 54.

Chlorolyne is a proparation of variable character, containing chloroform, ether, alcohol, oil of peppermut, hydrocyame acid, treacle, and morphine hydrochloride. Lobelin, capscum, belladonne, and extract of hydrocraresometimes added

Paragone Elizary is the popular name for the compound unotine of camplor, B P Various preparations, destintle of opaum, at o sold as "paragone substitute," &c., and if not dangerous in themselves, accustom ignorant passons to give and take large doses, which when repeated with genume paragone cause dangerous and even fatel effocts.

Distanton or Morsense and Denux—Liu cases of suspected possuming the detection of opinion is based, in addition to the recognition of its smell, on the extraction of morphia and meetine and in a sufficiently pure form to allow of the production of their characteristic reactions. The following is the usual mode of moreodura—

Observe if any smell of opium is apparent. If not, it may be stomech. Test a small quantity of the stained or filtered liquid with ferric elloride, and note if any red coloration (characteristic of mecome and) is modules.

Next ent up the stonach and any solid contents into small pieces, and radiace the whole to pulp by beating in a mortar. Mix the product with the liquid contents of the stormeth, and treat the whole with restribed spurt acadilated with sectic seed, in sufficient quantity to congulate the silminum 1 Keep the mixture warm for some time, with occasional agitation. Then filter or strain from the solid matter.

The filtrate is treated with basic accatas of lead as long as a precepitate is produced, when the hquid is boiled and allowed to cool. When cold it is again librered, and the precepitate weaked with cold water. The precepitate contains the meconic acid of any opum present. It should be weaked off the filter with water, and completely decemposed by passing a ranged stream of sulphurested hydrogen gas. The hquid is next filtered, and concentrated to a small bulk by evaporation at as low a temperature as possible It should then be placed in a procelum dish and tested with ferric childred, which will produce a purplish act coloration if meconic acid be present. It is necessary to distinguish enrefully between the coloration produced by meconic acid and the somewhat summar reactions given by thiocyanates and acctates. This may be effected with certainty as descuided on none 338.

A very useful indication of the amount of opium present may be obtained by comparing the depth of tant produced by ferric chloride with that obtained on treating a known quantity of opium in a similar way

The filtate from the lead precipitate will contain any morphine which may have been piesent. Separate the excess of lead by passing sulphuretted hydrogen for some time, filter, evaporate enthously nearly to dynose, add a little water and filter This filtrate will probably have a bitter taste if morphine (or other

Mocome and adheres very tensorously to albuminous matters, and hence the precipitate should be digested with strong sleohol, and the liquid strained and added to the main solution alkaloud) he present Transfer the solution to a stoppered separator, render the hopium dikulium varil ammous on (preferably) an alkalium bearbonate, and slake with hot amyhe alcohol without delay, as described on page 316. The amyhe alcohol solution is then separated, passed through a dry filter, and either at once evaporated to dryness, and the residue examined by the colour-test described on page 313 of seq. or it is slaken with a lattle didute hydroculioric acid, which is then separated and examined for morphime An estimate of the quantity of morphime present may be obtained from the intensity of colour produced by the iodic acid and ferrovandie tests (page 318).

Instead of treating the alcoholo extract of the maternal under examination with beats catestic of lead, as described in the foregoing process, the method may in some cases be abortened and rendered more deheate by ovaporating off the alcohol at a low temperature, taking up the residue with water, filtering, and altitude the filtrate with aliate sulphurue or hydrochloric soid, and squtaing with other? This temores mesome coul, though not perfectly, whilo phosphates and other interfering matters remain in the aqueous liquid, and if the ethered layer be espatiated, ovap-mited, and the residue treated with hot water, a solution is obtained, which after filtration may be very advantageously used for the application of the ferric chloride test. If preferred, the solution may be treated with lead acteta, and the mecome acad recovered from the filtered and washed precepitate by decomposing it with subhursteted hydrogen.

The positive detection of mecoma scal affords as perfect a proof of the presence of opum as does the recognition of mophicities?, and as the tests for and methods of separating meosure and from foreign materias are somewhat more satisfactory than the form form from the scale is more stable than the alkaloid, it occasionally happens that the acult may be solated and positively indistified, when morphine cannot be recognized with certainty (especially where plomannes may be pre-ent)? The detection of mecomic acid of course indicates the pre-existence of actual opium or some galennal preparation threeft, and not morphine or center the sales. Hence it sometimes canalies a useful distinction to be drawn as to the form in which he possion was taken

¹ After this treatment the aqueous liquid may be rendered alkaline with sodium brearbonate, and agristed with hot amyhe alcohol for the extraction of the morphine

² The author obtained satisfactory poof of the presence of meconic and in the stomachs of two children exhumed five months after death, whereas no positive conclusion could be formed as to the presence of morphine

It not unfi-quently happens, even in cases in which it is certain that opinin was the cause of death, that no trace of merpha or mercine acid can be found on analysis of the stomach or its contents. In other cases the poison has been detected with modernit facility a considerable time after death. The cause of these discrepant results is very obscure, but they are probably manily dependent on the opportunities which necronstances have given for the channitation or absorption of the poison before death. his ennued linear the first morphism of the poison before death has enued linear the first morphism of the poison before death. In section of the poison was not the cause of death. Attempts to extinct incepting from the blood and tissues have usually failed, but T. G. Wo in 1 sy has succeeded in nodating it from the brain, blood, liver, and urne of annuals possioned by it (Other News. km 7.9. 29).

In examining if in a for morphine, a considerable quantity of ures is lable to be taken up by the anythe alcohol. If the solution in this menstruum be evaporated and treated with cold water, a notable quantity of morphine is dissolved together with the unes. In the immute quantity present it may be extracted from the liquid by either (which does not dissolve urea), or preferably by a muxture of ether and acetic ether.

STRYCHNOS ALKALOIDS.

The vanous species of Stylectors, a genus of plants bringing to the order Logardaces, contain costum disclosis temericable for their miensely possionous properties. Of these, the only two which have been thoroughly investigated ane stryelning and brucing, the latter base being probably a dimethoxystypchilm.

Strychume and humane occur in the see ds of the Strychume mux commus, in combination with lactic and ignature axis! a hard base, igavarine, has been supposed to exist in mux commen, but the researches of W A S he arts one Glous Ohem See, xxxxx 433) have proved the supposed alkaloud to be morely a muxtare of staylename and brunen. The bar K of Shychine nume commus is also very possenous, and is sometimes termed "false angusture ata V". The extreme bittanese of the strychne bark, its twasted appearance, the unpossibility of separating it into bark, its twasted appearance, the unpossibility of separating it into that layers, and the blood-ted colonation produced on applying mitro and to the internal coat, are characters by which it is easy to distinguish it from true angusture bark

The seeds of Strychnos Iquation, commonly called "St Ignatius' beans," also contain staychnine and bracine, and are employed for the manufacture of the alkaloids, of which they are said to contain from $1\frac{1}{2}$ to 2 per cent ⁸

The leaves of Strychnos nux vomica are said to contain brucine but no strychnine 1

The decection of the root-bark of Strychnos Treuté or "deadly upas tree" of Java, evaporated to an extract, is the chief ingredient of the arrow-poison upas-ticute. It contains strychnine and lineine.

The deadly effects of Curare or Indian arrow-poison have been attributed to strychnine, but are now proved to be due to a distinct base, cui arine, which is described on page 388

Strychnine. Strychua CorHooNoOo,2

Strychnme exists, together with buene, in the seeds and heat of Strychnes mus vonue, in the seeds of S Lynatia, called "St Ignatia be ans," and in certain other plants of the same genus! It may be prepared from these sources by a method similar to that used for their assay (rage 385).

Strychnine occurs as a white powder, or in crystalline particles of variable appearance. The crystals are sometimes minute pearly

is Btychnine appears to have been found with centarity in five on are aposed of Sin_Advanced. Several of the genus contain nothine styphenic non-blucume 2 Accounts of the same of $1.8 \times ner$ (Zer. , vv. 773) the cirychinus of commerce has no talways the same components only gengeroscited in some matances by the formula $\mathbb{C}_{2}H_{2}N_{1}O_{2}$, and nothers by $\mathbb{C}_{2}H_{2}N_{1}O_{1}$, and believe the plant produces the alkinoid with a variable proportion of corbon, a supportion which has also been entertained by Schutzenbergen. Ke of to 4.9 by fracticatal procepitation of commencula strychnine with poissonin platnoschloride, obtained at first a salt containing 18 a per cent of Pk corresponding to a molecular weight of 347 for the alkinoid, while the precupitate missequently theory down contained 1985 per cent of plathnum, representing a neclecular weight of 347 for the alkinoid, while the precupitate missequently theory down contained 1985 per cent of plathnum, representing a commencul stypchnine probably contains how one strychnine $\mathcal{C}_{2}H_{2}N_{2}O_{1}$, and addition to the base of recognized composition

Nitr venues code or St Ignatus' basis are builed with disints sulphure, and till affet, then crubled, and the expressed hyund total valid sided lines in everes. The proceptate is offitteed off and boiled with abold of 6.8 species gravity, which disnover the alkalouis and deposits the strychmuse on cooling, the bruches mostly remaining in solution. The British Phon macapital distribute the powdrood cools shall be elastated with inthies abords, the spati-distilled off, and the solutions preceptated with acetate of Isod. From the listant tie halloudes are prompiated with another of Isod. From the lines to the listant to in alkalouis are prompiated with amounts and reducedoved is beling to official spart, the greater put of which is then distributed off. This rouseful liquid on cooling deposits the sky polinus, which is another with the contribution of the cooling about the sky polinus, which is another with the contribution of the cooling about the sky polinus, which is another with the contribution of the cooling about the sky polinus, which is another with the contribution of the cooling about the sky polinus, which is another with the cooling about the sky polinus, which is another with the cooling about the cooling

scales, like mnca, sometimes oelahedia, with a thombic base; but more commonly form large, four-sided primes. The neryslab vary much according to the solvent from which they are deposited. For their production on a miscoscopic scale it is best to let the alkaloul deposit goaldually by addition of an alkalia to the solution of one of its salts, o to expose the solution to ammoniace vapours (see page 36.1) Well-formed cryslas of strychmical relations of the from best produced by gradually adding water to the alsoholic solution of the from best.

Crystallised strychnine has an approximate specific gravity of 113 (T P Blunt)

Stychnue has no smell and is not delique/cent. On being heated it melts without decomposition at 265²-286⁸ C., and sublimes imperfectly. Its solutions exert a law-o-toknoty action on polarised light, have a marked slkaline reaction, and are strengly thitse.³

Strythnine is an exceedingly violent tetanic poison (page 372).

Strychnine is very sparingly soluble in cold water, requiring shout 8300 perts for its solution, but it dissolves in 2500 perts of boiling water It requires 207 marts of cold absolute alcohol for solution, and about 400 of whisky, 500 of spirit of 941 as gravity, and 2617 parts of 970 sp gravity. The limited strychnine in alcohol is utilised for its separation from brucine. which is readily soluble in the same bound. Strychnine is soluble m 8 to 10 parts of chlorofone, but dissolves very sparmoly in ether, requiring 1400 parts of the anhydrous menstraum, or about 1050 of ordinary commercial other Doubtless the physical condition of the alkaloid largely affects its solubility. Strychnine dissolves with facility in a mixture of equal measures of chloroform and ether-a fact often utilised for its extraction. It is soluble also in 140 parts of benzene, and is deposited on spontaneous evaporation in large brilliant octahedial erystals. In petroleum ether strychume is nearly insoluble, requiring, according to Wormley, about 12,500 parts for solution

Staychnine is not removed from its acidulated solutions by solution with any of the above immiscible solvoids, but, on the contrary, may be completely extracted from its solutions in them by shaking the hund with dilute sulchune acid.

Stayehmme is not sensibly soluble in solutions of the fixed caustic alkalies, but dissolves somewhat more readily in ammonia. In dilute acids it is readily soluble

¹ The bitterness of strychnine is said to be recognisable in a solution of right of a grain per gallon. The salts of strychnine are much less bitter than the free alkaloid. Stryohnmo dissolves without coloration in the strong numeral scales. It may be heated to 100° C. with strong sulphure and with-ut vasible change, and is often stated to be unalticed by such treatment. But the strychmine cannot be wholly recovered from the product, and C. Stochr (Be_{θ_i} , xviii. 3429) has shown that a sulphonic send is formed.

M on o bromstrychnine, $C_{23}H_{21}BrN_2O_2$, is obtained on the observation of attrohume hydrolyomide or hydrochloride, and them neutrating with ammonia The aqueous solution is alkaline and very bitter $(Arch \ Thorns, cas.xim. 318)$

SALTS OF STRYCHNINE

Stychnine is a strong bees, and forms salts which are usually cystallisable and soluble in water, yielding very bitter, exceedingly poisonous solutions. The salts of stychnine are mestly soluble in alcohol, but are ussoluble in ether, chloroform, benzene, petroleum sunts or amyte alcohol.

Strychnme may be titrated with accuracy by a standard muneral acid, using litimus on methyl-nange as an indicator. One or of deemormal acid conseponds to 00334 gramme of stychnine Strychnine, has no effect on phenolphthalem, and hence its salts react with filts indicator as if the acids were uncombuned.

The elephonate, perrocumule, mercu coldoude, phosphotungstate, and phosphomolybdate are among the most module sells of strychnina. All these forms are occasionally used for the isolation or estimation of the alkaloid. The high insolubility of the ferrocvande serves to seamnts the alkaloid from brunne.

The spanng solubility of the hydrocdule of stychnine is important, as the salt is liable to be thrown down in the form of crystalline needles from mixtures in which strychnine hydroliloide and a metallic rodde are dispend together. The hydroliloide is stated to be sumindry hable to separate out.

None of the salts of strychnine and any place in the British Pharmacopaia The subplate is official in the United States, and the nitrate in Germany The following table indicates the leading characters of the numerical salts of strychime

SIRTAININE ACCOUNTMENT ACTO, CAID, CALLAN, O., SO, H., is preduced in nearly theoretical amount when stychniar is heated to 100° with the requisite quantity of concentrated sulphian seed. The free acid as colourles, and very hittle colouble in water or alcohol. The anno nature masks is very soluble in note, but precipitated by alcohol, and the prinsimity colours, bearing, colours, bearing, colours, bearing, calcium, leaf, and copper sails form vay involuble precipitates. With funning sulphinus and at 150° a soluble dumly-horne used in formed.

			Proportion o		hility.
Salt	Formula	Appearance.	Strychnine	Gold Water	Boiling Water
Hydrochloride,	BHCI	Silky neodics	84 per cent	1 part in 50	
Hydrohromide,	BHBr	Prismatte needles	80 ,,	,, 32	
Hydriodide,	вні	Quadrangular needles, or white emics	723 ,,	Sparingly	
Nitrate, .	BHNOs	Silky neodies	84 ,,	1 part in 90	1 part in 3
Sulphate,	B ₂ H ₉ SO ₄ + 8 aqua (or 5 aqua)	Transparent quadratic octabulra	784 ,,	,, 42	,, 2
Acid sulphate,	BH ₃ SO ₄ +	Long, thin	714 ,,		14
Acetato,	2 Mas	Crystallises with difficulty	1	1 part in 95	湿

ANALYTICAL REACTIONS OF STRYCHNINE

- 1. On adding to a not too dilute solution of a soluble salt of strychum a fixed counts alkalı, alkalıne qarionate, ammonia, or hime-pikia, strychumine is himori down as a white precipitate moduble in oxcess of the precipitate. The precipitate moduble in oxcess of the precipitate The precipitate moduble becomes crystalline. The crystals have a characteristic microscopic appearance, being usually long, rectangular, well-defined puisar. They are well developed if a drop of a diute solution of a stychmne salt $(e\,p\,$, the acotate or sulphate) be placed on a shy of glass, and covered with a small beaker runsed with strong ammonia. After half an hour the beaker may be removed, the drop of liquid covered with a cruel of thun glass, and examined under the microscope of the solution contain extansions matter, it may be found difficult or unpossible to obtain a systals from it.
- 2. If skychune be hbented from the solution of one of its salts by one of the reagents mentoned above, and the liquid (with the suspended precentable) be at once shaken with an equal measure of chiloroforms, the althrolu is readily dissolved by the latter biquid, and may be obtained us solid state by separating the chloroform and evaporating it to dayness at a steam heat. The sgristion of the aqueous liquid with chloroform should be ropested if quarintistive results are desired. From aqueous liquid containing little solid matter, chloroform separates tolerably readily, but if, as often happens in practice, there be much extractive matter present, the complete separation of the chloroform requires many hours or even days. This meconvenience may be whelly avoided by substituting for pure chloroform a maxime of equal volumes of either and chloroform. This has a density of 1 II, and separates with facility from aqueous hands (owner pages 156 and 374). Experiments

by the author have shown that the solubility of strychnine in a mixture of equal measures of chloroform and ether is amply sufficient to cusure its separation from the aqueous liquid (Analust, v. 141)

3 A very useful precuptant for strychume in compiler organiliquids as a mine and solution of sodium phosphomolybalis (\$0 n n a n s h c n in * s n s g n n , negs | 136). On adding that sets of the solution of the alkaloud, the strychume will shown down as a yallowals white amorphous precuptate. The separation is complete even in very duthe bugins. Many alkalouds beades strychime give similer precuptates, and hence the suggests is merely of sevires for concentrating the strychime and purifying it from extraneous matters. The precuptate should be filtered off, washed with water containing the reagent, and the strychime separated by suspending the preciptate in water, adding summon, separated by suspending the preciptate in water, adding summon, and agrating with ether-chloroforn, as in test 2. The precipitate can, however, be directly excussed by the obour-reactions described on page 368.

4 Scheibler's reagent (page 136) precipitates strychnine from extremely dilute solution, and may be substituted (with

advantage) for the phosphomolybdic reagent.

5. Strychnine may also be separated from its tolerably concentrated neutral solutions by precipitation with chromate of potassum The test is best applied to a chloroform-residue obtained as described in 2 This should be dissolved in dilute acetic acid, the liquid filtered, if necessary, and evaporated to dryness at 100° The resultant acetate of strychnine is dissolved in a little cold water, and neutral chromate of potassium is added to the solution. Strychnine chromate, (CorHoseNoOo), HoCrO, is thrown down as a reddish or yellowish brown precipitate, soluble in boiling water (1 in 171) and ie-deposited on cooling in orange- or lemonyellow needles and plates The precipitate is very slightly soluble in cold water (1 in 470), a fact which enables strychnine to be separated from brucing, the chromate of which is more soluble Potassium highromate throws down from solutions of strychime. not too dilute, an anh v droch i om at e of the formula B.H.Cr.O. as a crystalline precipitate, in which octahedra and bush-like groups are the most prominent microscopic forms. The precipitate is not soluble in excess of the reagent or in very dilute acids, and its formation is much facilitated by stirring. It dissolves in 1800 parts of cold and about 240 parts of boiling water, and is rapidly affected by exposure to light. The chromates of strychnine give the characteristic violet exidation-product directly on treatment! with strong sulphuric acid as described in paragraph 8, or the alkaloid may be obtained in a free state by suspending the precipitate in water, adding ammonia, and agitating with ether-chloroform, as in 2.

- 6 With induced potassium, nothic stychime gives a reddishown pracepitate, even in extremely dulute solutions (1 100,000). Mayor's reagent also precipitates strychime from very dilute solutions (T 150,000), and is recommended by G. F. Schacht for its determination.
- 7 Strychume forms a combination with jodine analogous to. and having similar optical properties with, herepathite The following is the best method of utilising the reaction for the detection of stavehnine. On a microscope-slide place a very small drop of an alcoholic solution of rodine, and allow it to evaporate Directly it is dry add a drop of a solution of strychnine, made by dissolving the alkaloid in dilute accide acid and adding a drop of sulphung acid. Add also a drop of rectified spirit, and allow the mixture to evaporate spontaneously. On examining the residuo under the microscope with a Nicol's prism and selenite, but using no analyser, characteristic crystalline structures will be observed These may take the form of small cucular tufts of very fine black ncedles, of minute dots of a more or less triangular form, exhibiting vellow, pink, and green tints, large triangular crystals of a yellow or green colour, composed of three parts radiating from a centre, numerous solid macled prisms, occasionally showing complementary tints, or solid resettes of four five, and six sided prisms. In all cases it is desirable to compare the results with those obtained from a minute quantity of strychnine treated in precisely the same manner. The mode of operation may be varied considerably, provided that the essential conditions of simultaneous presence of alcohol, sulphune acid, acctic, acid, free nodine, and a trace of staychnine be duly observed. The test is said to be sensitive to 1-2500 of a grain of strychnine
- 8 When poissoum ferracyande is added to the solution of a sit of strychinine, the ferrocryande of the base, \$M,FeCy_2+3H_0,\$ is prequitated as a white crystalline powder with a shade of yallow, only very spanningly soluble in cold vater. The observation, which is due to Beckurts, has been utilised by Dunstan and Short (Fen-Book Flearing, 1883, page 468) for the determination of strychinine and its separation from lucinous, the fearory-anile of which is readily soluble. A quantity, not exceeding 0.2 gramme, of the muxed alkalonds is dissolved in about 10 cc of water containing 5 per cent by measure of strong sulpurae each the solution diluted with water to about 175 cc, and them made up to 200 cc, with a 5 per cent, aqueous solution of bostssum ferovanied. The hound is strand occasionally during

six hours, and is then filtered off and washed with water acidulated with 10 of sulphune acid, till the washings are free from bitterness As the procuptate is hable to alteration on diving. 1 it should be washed off the filter with strong ammonia and extracted by agitation with chloroform After separating the chloroform solution and washing it with water, the strychime may be titrated by standard acid and methyl-orange, or the chloroform may be evaporated to dryness and the residual alkaloid weighed. Some alcohol should be added towards the end of the evaporation to prevent the violent decrepitation which otherwise ensues.2 From the filtrate from the ferrocyanide precipitate the brucine may be precipitated by ammonia and extracted by chloroform Schweissinger (Archiv des Pharm, [3], xn 579, 609) states that he had not found the ferrocyanide method to effect a perfect separation of strychnine and brucine. He found strychnine ferrocyanide to be perfectly mesoluble in water acidulated with sulphuric acid, but the brucine salt was not completely soluble, and was precipitated more or less perfectly after a time. Hence the strychnine was always estimated too high and the biucine too low, the error largely dependme on the time allowed and the concentration of the hould

When the precipitation of the strychime as feirogrands is effected in should stongly send with hydrochlore send, the salt thrown down is insoluble in cold water and alcohol, has a blunch shade, and is an acid feir long standamy Bright Strych No minder precipitate is obtained with butchine except in lightly concentrated solutions, on after long standing. Holst and Beckurts (Arch. Phan m. [3], xxv 313) have based on time feet the following columetric method of determining strychime. Al per cent solution of the alkaloids is strongly accidiated with hydrochloric scali, and a standard solution of potassium ferrocymind solution added until a filtered potion of the highid gives a blue stan with pope moistened with ferric chloride 294 per of feirocymide represent 334 of strychime. The following results were obtained:

¹ According to Beckuits, upon exposure to air staychnine ferioryanide turns yellow, and is eventually completely decomposed with formation of strychnine ferioryanide and a new base which can be extracted with alcohol, celled exect yellome, Ca.La.N.O.

² Dunstan and Short state that this behaviour is characteristic of pure staychine, a minute admixtate of brucine preventing it and ceasing the alkaloid to have a fused appearance

- 9 On teating a cold solution of stychnine in concentrated sinch purple-blue coloration is developed. This changes more or less rapidly through jumple and ermison to a bright cherry-red tirt, which is somewhat persentent. The rapidity of the change of colour is largely dependent on the nature and quantity of the conclusing agent employed. Valuous substances have been recommended for the purpose. The following are the most notable.—
- (a) Polassium belromats This is a favourite oxidieng agent with many openitors, but in the expenseor of the author is one of the least reliable reagents for the purpose, set he change of colour is very rapid and the giene chomium compound resulting from the reaction tends to mark the colonation due to the strebune
- A useful way of employing bichromate is to piecipitate the strychime by means of is (as in 5), and apply sulphime acid to the precipitate. This plan has the great advantage of separating brugue, the presence of which is objectionable.
- (b) Polassium permanginate, originally isocommended by Guy, gives the reaction with great distinctness, but the rotation of tints is very input, and the reagont itself is apt to gave a crimison colour with subhuric acid.
- (c) Potassium ferricianide, a reagent proposed by E Davy, gives exceedingly good results The change from blue to crimson and red is very rapid
- (d) Lead dioxide (PbO₂) This oxidising agent, suggested by Marchind, acts immakally well, but the puce colour natural to it is apt to distract the attention from the reaction to be looked for
- ack, employed an moderate quantity and in the finally powdered state, is the one to which the author gives professore. The play of colours is remarkably well-developed, and the change of tart very enabled
- PR Mandelin recommends a solution of 1 gramms of aummonum vanadate in 100 cc of sulphure and as a reagent which will keep nuchanged, and which gives the colour-reaction with great distinctness
- (f) Cooseer could (Cc,O,) has been highly recommended as the exclusing agent by S D Hinsdale. It has the advantage of being light in colour, and giving a colouless reductionproduct.

The oxidation-test for strychnine is usually performed in practice

on the residues left by evaporating to drynoss the ether-olloroform with which an alkalms solution of the alkalood has been egitated. The test may, however, be directly applied to the chromate or phosphomolybelate of strychmine (see reactions 4 and 5). The following mode of operating is best calculated to ensure delicacy and accuracy "

The solution of the strychnine in ether-chloroform should be evaporated in a porcelain dish or crucible. If the quantity of strychnine to be sought for is likely to be very small, the dish should be immersed in hot water, and the solution of the alkaloid allowed to fall slowly into it from a burette or pipette, so that each drop may almost completely evaporate before another arrives In this manner the strychnine-residue may readily be confined to a very small area, and the after-reactions thus rendered proportronately delicate When quite dry and cold the residue should be treated with two or three drops of pure concentrated sulphuric acid, which should be thoroughly incorporated with it by means of a glass rod. The mixture should then be allowed to stand for five minutes in order to note if any colour is produced Salicin and certain other bodies will cause a red coloration, while some may be more or less charred. If any marked coloration is produced, the dish should be gently heated (not to the boilingpoint of water) for half an hour, the contents diluted with water, filtered, made alkaline with ammonia, agitated with a mixture of ether and chloroform (as in test 2), and the strychnine recovered by evaporating the solvent. The residue is then again treated with a drop or two of sulphuric acid

The oxidesing agent, which should be, by preference, manganess or lead dioxide, is then added to the sulphune and by dipping a glass rod moistened with the latter into the powdered selid. A modester quantity only should be used, so as not to obscure the reaction by excess of blackness. On strring the drop of stryction will become developed. In a minute or so it will be distinctly to a construction of the contract of the contra

There are but very few substances which at all simulate the reaction of strychime when treated with sulphuic and and an

¹ The oxidation-reaction has been applied by Davies and Sohmidt to the approximate determination of the strychnine in Easton's Sylup (Year-Book Flasm, 1883, page 671).

oxidising agent, and few indeed of these that are dissolved together with strychims on agricting the alkaline solution with ether-schoofform. Saliein, santonin, piperine, solanine, extens opium bases, cod-liver oil, indication resins give colours with subplanue and alone, but they are extracted from and solutions by ether and chlorofform, and certain of them may also be got rid of by gently heating the laquid as already described Aniline gives no colour with subplanue seed alone, but obcured products are formed on treating the solution with an oxidising segent. These cannot be mistaken for the oxidation-products from strychime, for the oxide of this sentincy different, commencing, in the case of aniline, with a green, changing to a very persistent blue, and ultimately becoming black. Colory in his energy was a very similar rescion to strychime, but is readily extracted by antitum the seadulisted solution with beause or sther.

It is always desirable to purify the shyelmine by extracting it time an alkalmen beput by againston with ether-chloroform (see page 364), but the exhibiton-reaction is readily obtained even in presence of conaderable quantities of certain foreign matters. Thus ost-meal, tartai-emesis, and dextain do not materially interface with reaction when the quantity of strychimue is considerable. Some extractive matters, sugar, and nitrates wholly prevent the application of the colour-test, and thereof the absence of strychimum sunear be assumed till the test has been applied to an ether-chloroform results.

Quinine, einchonine, and veratrine may be found with stychnum in the sthe-chloroform seather, but do not metall proportion with the application of the test. Morphine in small proportion does not intelled, and the presence of any larger quintity in tances is excluded by its limited solubility in the ether-ablosoform in small proportions by using excession on inturious informa-

on the condition-less for strychmus, but when much is present it interferes in a marked manner. Hence it is safest to separate the strychmus first of all as chromattee or ferroryande, as described in reactions 5 and 7, or a strong solution of a sait of the allacland can be treated with a ray feeded excess of amounts, when the stychmus will be presentated and the brucace will remain in solution. If a maxture of brucane and strychmus be treated with chlorus—water, the former base dissolves as a lacklorobust on the structure of brucane and strychmus be treated with chlorus—water, the former base dissolves as a lacklorobust on the structure of the sum of the structure of the alkaloids, rathes than to the use of a nutrileal natural sources of the alkaloids, rathes than to the use of a nutrileal

salt of strychnine Commercial strychnine and its salts often contun traces of brucine, but not sufficient to interfere at all with the application of the oxidation-tost.

Curaline, the active principle of the Indian arrow-poison, gives a series of coloured oxidation-products oxacily like those of strychnine, but not being sensibly soluble in chloroform it is not hable to be found in the chloroform-residue (see page 359)

A p to main a has been described by \tilde{G} Am thor (Ohesenset, x, x, x, x, x), which gives a blue colour with the condationative less persistent and pure than that produced by strychime. It is less inter and less presences to fings than strychime, is dissolved endily by amplic alcohol but only slightly by either from alkaline solutions, and gives an among house chromate, picrate, ferrocymide, and foreign and proves an among house chromate, picrate, ferrocymide, and foreign and gives an among house chromate, picrate, ferrocymide, and

Many of the above sources of fallacy or confusion may be wholly avoided by performing the oxidation-test in a manner suggested by H. Lethely, which comusts of employing electrolytic oxygen instead of either of the exiding ngestin mentioned on page 368. The solution of the either-chloroform residue in a drop or two of strong sulphure and is placed in a unp-shaped depression in a pace of platinum foil. The foil is connected with the platinum plate of a single Grove's cell, and a platinum were connected with the zinc plate of the battery. Immediately that the ond of the platinum were is dipped into the doctor of and, the voide colour of the oxidation-product will flash out, and on removing the wire from the lound the time will remain.

8 A colour-reaction of strychnine with chloride of zinc is described on page 145.

 $U_{\rm e}$ 1f solid siryolmane be dissolved in a drop of dilute intro cascal, the luquid gently heated, and a minute particel of potassical characteristics and the siryolman characteristic control of the variable of variable o

10 A reagent prepared by adding sufficient strong hydrochloric acid to a weak solution of potassium chlorate to render it bright

¹ The reaction may be rendered still more delicate by placing the drop of hound at the bottom of a porcelain ciucible, and momentarily immersing in the liquid two platinum wises connected respectively with the zinc and platinum plates of the battery.

yellow, and then sufficient water to make it a very pale yellow, gives with a solution of strychnine in hydrochloric acida fine red colour, destroyed by excess and restored by boiling Erucine gives a violet coloration (C L. Blo x s.m., loc. cit.)

11. An exceedingly delicate test for strychnine is the physiclogical one of Marshall Hall A freshly-caught frog the smaller the botter, is the best subject for the experiment. The skin of the back should be raised with a pair of forceps, and a small sht made with a pair of scissors. Into the opening, the suspected liquid, as concentrated as possible, should be injected by means of a small pipette. The first symptom observed will be a difficulty in breathing, which gradually increases till the animal appears to gasp for breath A slight tremor will be observed extending over the whole body, but specially noticeable in the hand legs. The frog sometimes remains perfectly quiet, but in other cases takes energetic and convulsive leaps. It should be placed under a beaker or bell-glass for easier observation. The characteristic tetanic convulsions next make their appearance. They are intermittent, the pupils being dilated during the spasms and contracted in the intervals. The convulsions may be induced by touching the frog, clapping the hands, or knocking on the table

The physiological test is much reduced in practical value by the difficulty in obtaining young animals for experiment. On the whole it is deadedly less certain and characteristic than the chemical reactions, and in no case should be implicitly relied on unless confirmed by the results of the oxidation-test.

Toxicology of Strygining

Owing to the violently poisonous character of strychnine, and the ease with which its preparations (under the disguise of "vermin-killers," &c.) may be obtained by the public, cases of death from its effects are very numerous.

The symptoms of possessing by stripolanus usually commence with a bitter taste, followed by a feeling of suffication. The characteristic tetanic convulsions, often accompanied by opisthotonos, then come on, gradually becoming more frequent? Vomiting is not common. Lockjaw is a constant symptom. Consciousness, as

Methyl-strychnine produces a paralysing effect more allied to that due to curare than to the tetanising effect of strychnine.

¹ In the author's own experence of the examination of posioned animals, oxtending over amany years and to a great number of cases, strychnine has been a found more frequently than all other kinds of poison taken together. He has a most with it in several cases of murder of human burge, the criminals subsequently undergoing capital purashment, and in numerous cases of sincels and death by musclevinters, unboulding carless dispensarily by a qualified medical man.

a rule_is retained till the last accompanied by a lively terror of the rapidly-recurring and agonising fits. Death usually ensues within a few hours, but in raic cases life has been prolonged for several days. The general time is from thirty to musty minutes!

From $\frac{1}{2}$ to $\frac{1}{47}$ of a gram is the usual medicand done of stypching, but I may be increased in the case of a person accustomed to it. One-exith of a gram is usually distinctly dangerous. One grain may be seguided as the average fatal does for an adult, and death has been known to occur from $\frac{1}{4}$ gram. Much larger doses have been recovered from 2

// Hypodermic injections of strychine have been very successfully employed as an antidote in cases of snake bite.8

The post-mortem appearances of posoning by strychime are not very striking or characterists. Rigidity of the muscles is usually prolonged, but if death occur in one of the intervals between the fig. no rigidity will be observed. The heart is usually, but not always, full of blood, especially on the right side. The stometh usually appears normal, but sometimes intensity concessed. The

In a case within the author's expension, in which medicine containing a possionist does of strychinine was taken, the victim, a young woman, immodiately suisl out that she was poincid, and died in reslive minutes. Analysis of the remainder of the medicine showed the piecene of rable more than one nor suit of strychinine in such does, and the amount of poinci necessary different the viscous accorder formatically closely with this result.

¹ The most snoesaful antidote for strychinion as the presistent inhalignout. Algorithms of the presistent inhalignout. Algorithms of the presistent inhalignout. Algorithms of 20 grants, has proved highly efficacions on several occasions, in acute instances the cramps beang wholly prevented, while, on the other hand, on nanoritie action of, the antidote was in "" I "" has proved in a nanoritie action of, the antidote was in "" I "" has proved in a nanoritie action of, the antidote was in "" I "" has proved in a nanoritie action of, the antidote was in "" I "" has proved in the prevention of the antidote that the province of the prevention of the antidote for stream of the prevention of

⁸ The strybhume as used as untate un 240 parts of water (~6 gnama to the conceop maxed with a little glycorm. Twenty mumma should be unpreted every 10 to 20 manutes until all the snake-posons symptoms have thespeared and sight muscals; against are observed. A gener ur more of strybhume may las required in the course of a few bours. On the substitute of the size of the state of the size of the s

⁴ In a case in this author's experience, the stomach presented such an appearance as to suggest the presence of an assence or other irritant posson, but no minical posson could be detected. That death was due to administration of a vermun-killer containing strychnine was subsequently fully proved by analysis and admitted by the munderer.

most characteristic appearance is the intense congestion of the brain and spinal cord, often accompanied with considerable effusion of blood

For the detection of strychnine in the dead body, the following method should be used, the portions of the body operated upon being chosen according to the manner in which the poison is likely to have been administered. Thus it is of no use to search in the stomach or intestines for strychnine injected hypodermically. If the poison has undergone absorption, it will most probably be met with in the liver, but all parts supplied with blood and most of the secretions may contain small quantities of the poison. In extreme cases, it is desirable to operate on very considerable quantities of material, as death may be caused by so small a quantity of strychnine that the noison may be altogether missed

if this precaution be not taken

The portions of the body to be tested for strychning should be cut into small fragments with a pair of seissors, and then further reduced by bruising in a mortar. The product is then treated with rectified spirit, mixed with about 1 part in 20 of acctic This coagulates the albummoids, while allowing of the complete solution of the strychnine After a few hours the liquid should be strained through mushin, and the clarified filtrate passed through a paper filter. The clear liquid is noxt evaporated nearly to dryness, diluted with water, and again filtered. The filtrate is once more evaporated to dryness, and the residue thoroughly extracted with strong, and preferably absolute, alcohol The liquid is filtered, the alcohol removed by evaporation, and a small quantity of water added. The solution is placed in a tapped separator, diluted to about 20 c.c. with water, and a few drops of hydrochloric or dilute sulphunc acid added. An equal measure-of ether is next added, and the whole well shaken. On standing a few minutes, the ether will separate on the surface, when the aqueous hauld should be withdrawn through the tap, and the other then run off into a separate vessel 1 The aqueous liquid is then returned to the separator, and about 30 c.c. of a mixture of equal volumes of ether and chloroform added Enough ammonia to render the liquid distinctly alkaline is next added, and then the whole immediately shaken thoroughly for about a minute. On coming to lest. the aqueous liquid will tend to separate from the mixed chloro-

¹ This probusinary treatment of the acidulated solution with ether is very important. It effects a separation of glucosides, traces of fat, essential oils, and other matters which otherwise would contaminate the strychnine. In some cases it is desirable to repeat the agitation with a mixture of equal measures of chloroform and ether.

form and ether, which has a density of about 11 If telenably free 'from extractive matter, it will float on the surface of the other-chloroform, but if largely charged with sugar or other soluble matter, it, may be equally dense with the solvent, or even collect at the lower part of the separator If, from the presence of extractive matters or for other reason, the liquids do not readily separate, water or ammona should be added, so as to reduce the density of the aqueous liquid. An alternative, and perhaps proferable plan, is the gradual addition of ether, with cautious agration, till the solvont separates readily at the surface of the aqueous liquid.

When the division of the contents of the bulb into two layers is complete, the strata are separated from each other by means of the tap If quantitative results are required, it may be desirable to agitate the aqueous haund with a fresh quantity of ether-chloroform The solution of the alkaloid in the ether-chloroform is passed through a small paper filter, if necessary, and then evaporated to dryness at a steam-heat in the mannel described on page 369 The residue obtained may then be examined for strychnine by the tests given on page 364 et seq. If strychnine be present, the solution of the residue in alcohol will have a marked and persistent bitter taste, especially noticeable at the back of the tongue. The most delicate and characteristic chemical reaction of strychnine is the exidation-test described on page 368 Reactions 5, 7, and 8, and the production of crystals of stryclinine as described in 1, are also valuable as confirmatory tests, and should never be omitted if the material at disposal be sufficient for their performance. The bitter taste, however, in conjunction with a distinct reaction by the characteristic oxidation-test, may usually be regarded as ample proof of the presence of strychnine, provided the absence of interfering substances has been ensured by the previous treatment. The ptomaine, stated by C. Amthor (page 371) to give a colour-reaction simulating that of stryclinine, can only be present when putrefaction has taken place, and its formation must be very rare, or it would have been met with in the numerous cases in which no alkaloidal substance has been detected

Blood should be examined for strychnine by diluting it with an equal bulk of water, adding a little acetic acid, boiling for a

¹ This alternative is preferable to the addition of ablosform, which, if used in too large a proportion, will only separate from the dense aqueous liquid with extreme difficulty. The advantage of employing a mixture of other and chlosform, rather than either solvent singly, has been pointed out by the author (Asnaba, vi. 141), though it use did not originate with him.

short time, filtering, and evaporating the filtrate nearly to dryness The residue is taken up with alcohol, and the solution treated as already described.

From urine, strychnine may be directly extracted by agitating

the fluid with ammonia and ether-chloroform,

Dialysis through parchment-paper is an efficient and occasionally a convenient means of separating strychnine from organic matter. The finely-divided tissue should be suspended in water, to which some alcohol and acetic acid have been added. Distilled water should be used on the other side of the membrane. and changed at intervals of twelve hours. After thirty-six to forty-eight hours the dialysate may be evaporated to dryness, and treated with alcohol, &c, as described on page 374

It has not unfrequently happened that a post-mortem analysis has failed to detect strychnine in corpses almost certainly containing it. This result has probably been due in most cases to the use of defective methods of analysis, or to the search being restricted to too small quantities of material or to wrong parts of the body Occasionally, failure has probably been due to an elimination of the poison during life, especially in cases in which death has resulted from a minimum dose. Strychnine does not undergo decomposition in the dead body, and has been detected several years after death.1 Hence, if elimination has not occurred prior to death, strychnine ought to be found by the toxicologist.

PREPARATIONS OF STRYCHNINE

The only preparation of strychnine recognised in the British Pharmacopæia is a solution of the hydrochloride, which, as met with in commerce, is not so constant in strength as is desirable.

Easton's Syrup is a widely-used remedy, consisting of a syrup of the phosphates of iron, gumne, and strychnine Its omission from the British Pharmacopæia is lamentable, and results in considerable variation in the composition of the preparations sold under its name According to Squire (Companion to the British Pharmacopesa), when prepared according to Dr Easton's formula, the syrup contains "about I grain phosphate of iron, I grain phosphate of quinine, and all grain phosphate of strychnine in each fluid drachm "

¹ The author has had no difficulty in detecting strychnine in a stomach presorved in spirit for six years. A portion of the untreated etomach and liver from the same person (who picked up in a field and ate an egg poisoned with strychnine) was kept in a jar, the mouth of which was closed by a bag containing wood-sharcoal On opening the jar after six years, the whole of the contents were found to have disappeared, with the exception of a small quantity of dust, in which abundance of strychnine was detected,

The following is the range of variation observed by Davies and Schmidt (Year-Book Pharm., 1883, page 575) in ten samples of Easton's Syrup of commerce.—

	Squire's	B P Committee's		Found		,
	Formula.	Formula	Highest.	Lowest	Average.	
Quinine phosphate, Qu _s (H ₂ PO ₂) ₀	6 87	6.0	7:68	1.67	5 00	Grains per fluid or.
Forrous phosphate.	5 80	8-0	12 82	0 97	6 91	110 02.
Free phosphoric sold,	88 03	500	49 24	19 86	84 88	13
Strychmine,	10	10	80	0.6	10 to 114	Grains por
Spoolfic gravity,			1 881	1 288	1 298	* nuit or

The following analyses of commercial Easton's syrup have been published by J. G. Wilson (Phaim Jour, [3], xix. 753) —

			A	В	c	D	E	
Quinine phosphute,	-		6 75	6 75	5 25	4 25	2:00	1
Ferrous phosphate,			71	7.5	64	50	50	24
Phosphorio sold,			47 0	45 0	48 0	81.0	26 0	Grain
Stryolmine,		٠	0 25	0 25	0.25	0 20	0 10	10-

In analysing Easton's syrup the *iron* may be determined by evaposating 5 cc of the preparation, igniting the residue, discolving the sah in hydrochloric acid, and titrating the iron with standard bichromate solution after reducing it to the ferrous state.

The free *phosphoric acid* may be determined by titration of 10 cc with methyl-orange and semi-normal caustic sods. The neutral point is attained when NAFLPO. is formed.

point is attained when NaH₂PO, is formed.

The alkalouds are determined by diluting 10 oc of the syrup with twice its messure of water, adding some citic acid and oxcess of ammonia, and agitating twice with ether-folloroform. From the weight of the residue left on evaporating the solution, a deduction of 0 0057 gamme should be made for the strychime present, the remainder being regarded as quietims. An actual aspention can be made by prompitating the strychime from a strongly and solution by profession ferrogrande, as described on page 367

Another method of separating the strychnine and quinine of

¹ From the aqueous liquid the total phosphonic acid may be thrown down by magnesia mixture

Easton's syrup is to dissolve the ether-obloroform residue obtained as above in about 10 c of water estimated with a few drops of sulphurne need. The solution is neutralised by animonia and mixed with excesse of animonium oxalate. After standing twenty-four hours, the precipitated oxalate of quanne is filtered off, the mother-house removed by gentle pressure, and the precipitate washed once with a little cold water. It is then direct al 10° and weighed 1 Ke weight, multiplied by 878, gives the quanne in the quantity of the sample operated on The filtate and weak-water are then treated with ammonia, shaken with ether-chloroform, and the discovered alkadod (consisting of strychime, any amorphous alkadod, and a more trace of quinney should be next twice treated with 3 cc. of washed ether, which dissolves it

For the determination of the small proportion of stydenus contained in Eastorie syrup, Davies and Schmidt resonamend the following colormetric process devised by O. Ecken is term. The stakeholal resulte from 10 c. of syrup was dissolved in 31 25 c. of water acidilated with 1 c. of normal sulphune acid, and 5 drops of the resultant solution were added to 4 c. of concentrated sulphure and inited yellow with potassum behometa. The colour produced after standing five numities was then compacted with the colour produced by known quantities of a very dultie solution of strychinus of known strength, in this asme sulphune acid coloured with bichromate. For quantitative purposes the method leaves much to be desired.

Easton's syrup is hable to give a deposit which sometimes contains quinine, and in other cases appears to be simply ferric phosphate. The tendency to deposit is often prevented by addition of a small quantity of hydrochloric acid.

Versum-Aillers. An inquiry into the composition of various commencial vernim-killers containing strychnine was made by the author in 1889 (Year-Book Phan m, 1889, page 434). The results showed them to consist of a mixture of strychnine with nece or wheat-shach, usually more or less coloured. Ultanamene was the most usual colouring agent, but prussan blue was met with in four picquartantons out of seventeen examined, in one case the powder containing both ultramarine and prussan blue. Carmine

¹ The mode of operating described in the text is due to B. W. D war a It would probably be better to wash the promptate produced by ammonium oxalate, and then extract the quinino in the free state by agritating the precipitate with ammonia and other.

was the colouring-matter of two preparations and soot of one,¹
In one instance, no colouring-matter whatever was present,²

Ultramerine is leadily recognised in a vermin-killer by the peculiar shade of blue it communicates to the powder, and by the colour being wholly destroyed by agitation with dilute acid. If a little of the powder be placed on a silver com and moistened with dilute acid, a brown stain will be produced on the coin by the sulphuretted hydrogen liberated from the ultramamne. Ultramarine tetains its blue colour after ignition, whereas prussian blue leaves a brownish residue of oxide of iron, and indigo is more or less perfectly consumed, according to its purity. A decidedly ferruginous ash is left by some specimens of indigo. Prussian blue and indigo are unaffected by dilute hydrochloric acid. If the residue left after heating the powder with dilute hydrochloric acid be washed and treated with caustic soda solution, it will be unaffected if composed of indigo, but prussian blue will be turned brown, and the filtered liquid will contain a ferrocyamde, and hence will yield a blue or green precipitate or coloration when it is acidulated with hydrochloric acid and ferric chloride added

The colour of a vermu-hiller should not merely serve as a dangur-signal, but be so chosen as to facilitate its detection in cases where it has been used for the purpose of suiede or number. In a case in which the author was conceined, a murderer would probably have seesped conviction but for the detection of the blue colouring-matter in the stomach of his victim, which served to connect him with the administration of the nossus ⁹

¹ This preparation consisted of strychnine, 5°8 per cent, native builtin carbonate, 45°0 per cent; and wheat-flour and soot, 49°2 per cent. The object of the combination is not evident.

Such a colourless preparation is highly dangerous. Toothing provides no so generally coloured pink that they are not unfrequently saked for as "mak providers," and gray provides are equally common. The bine colouring-matries present considerable advantages over such guencies as soot and curranno, since no food, drink, or medicane has naturally a blue colour, and hence the unit at once attracts attention.

2 Soot is manitable for colouring remus-billers, as the sitentification of multise particles of cubon in difficult or impossible when muzed with food of the blue colouring matters practically available, ultramarane is to result discrept oily duties early and present pure, though the the stratage of being readily detected, and of being undestroyed by sginton. Plusian blue is unseffected by acids, and one two vay readily affected by district all the layers. Independent of the colouring agents, and an early and a not affected by acids, care also decolorised by all that the strategy and a not affected by district and in the one of the colouring agents, and in all decolorised by all that me relating agents in munte quantity it is less easily prospiled that purposes allow. A matter of the three pagements would be

The toxacity of veanm-killers varies within wide limits Of the samples examined by the author, the weight of strychnine contained in a packet of the powder varied from 0.60 to 4.18. grains, the relial price in each case being 3d. The proportion of strychning named from 4.2 to 4.18 nor cent².

Strychnme can be determined in vernm-killers by exhausing, a known weight of the dry powder with chicorform or beausing, and weighing the alkaloidal reaction for on evaporating the solvent The inschible portion must be examined by the taste and exidation-test, to ensure complete extinction and the absence of a still of step-thine inschible in the solvent used. An alternative, and in many respects preferable, method is to treat the verniller with cold water senduisted with sector acid, until the readual powder has no britter teste, and gives no coloration by the exulation-test. The solution is then treated with excess of aumonia, and the strychnine extincted by ether-chloroform, which is senanted, evanowised to drivens, and the readule woulded.

Of vermin-killers containing strychnine, Battle's preparation is the best known, and most extensively used. The suicides

preferable to any one or two of them. "The most suitable pigment for colourmg vermin-killers would be chrome green (Or2O2) In it we have a bright green pigment of high coloning power, quite insoluble in water and dilute acid and alkaline liquids. It is wholly permanent under all imaginable conditions, and is not affected by ignition. Chromium is not a natural conetitiont of the body, is not used internally as a medicine, and is not liable to be present accordentally, even in traces, in any beverage or miticle of food. It can be detected and determined with case and certainty, even when present in very minute quantity Owing to its meolibility, oxide of chromium would remain wholly unabsorbed if taken internally. Hence, if it were added to preparations of strychnine, &c., in a definite and invariable proportion, an estimate of the minimum amount of poison taken by a deceased person could be arrived at by determining the amount of chromium contained in the alimentary canal, even though the poison itself had been wholly absorbed or decomposed, and this could be effected with equal case and certainty after prolonged inhumation, or even after cremation of the body "-A. H Allen (Year-Book Pharm., 1889, page 489).

11 does not follow that the vermu-cluller which contains the greatest weight on the largest reportion of strydmum as the best for its process. Clearly, pure strydmum as would be meladent, and have the color stellar companible with the attractive and appetency character. To effect this, the letter cases of the strydmine should be such color of the strydmine should be masked only preside, and a straight colorest the strydmine should be masked as the stream of the strydmine should be masked as straight colorest the strydmine should be masked as the stream of the strydmine should be made to a straight colorest the strydmine should be made to a straight colorest the strydmine should be made to straight and to straight the straight and out of same. Most of the vermu-childra examined by the author have been doorlies.

due to it amount to many scores, and probably to hundreds. The colouring-matter of Battle's vermin-killer appears to have been uniformly prusuan blue, but the following table shows that the composition ascribed to the proparation has varied in other respects at different periods:—

Authority	A Swaine Taylor	A J Bernays	T Steven-	A H. Allen	Tardieu	Woodman and Tidy
Date,	1862.	1876.	1883.	1889		
Price of packet,	8đ.	84	66	64		
Weight of powder,	13 grains	15 grains	25 grains	21 8 grains	20 grains	
Colouring-matter,	Prussian	Prussian	Prussian	Prussian	Prussan	Prosessan
Starchy matter, .	blue Flour	blue Wheat-	blue	Wheat	Potato-	blue Flour
Strychnine, grains,	0.78	flour 16	28	flour 2 4	starch 25	
Strychnine, per	58	10-7	100	11-2	77	25 0

The nert matter of vermin-killers usually consists of rice-starch, though in some cases wheat-flour, and occasionally oatmeal, is substituted. In one instance, the author found both rice and wheat starch, the powder being coloured with carmine

Brucine. Brucia. C28H26N2O4; or C22H18(OCH8)2

Brucine, occurs in association with strychnine in nux vonuca, St Ignatius' beans, and false angusturs bark (page 360). The leaves of strychnos nux vonuca are stated to contain brucine but no strychnine.

In chemical constitution, brucine appears to be a dimethoxystrychnine 2

Brucia occurs as a bitter, white, odourless, ciystalline or amorphous powder, or in groups of very deheate needles or foursided prisms, containing 15 45 per cent of water ($=C_{28}H_{20}N_2O_1+$

1 Woodman and Tidy state that sugar is a constituent of Battle's vermin-killer. This was certainly not the case in 1899. The proportion of strychnine (23 per cent) given by Woodman and Tidy is laigely in excess of that found by other observers.

² He uses n finds that both strychume and breuze yield by oxidation with chromous end matures a body contauming C₀H₀H₀A₀, and show that the difference between the two alkalosis must be sought in the residues, C_HE₄ and C₁H₀O₇ respectively, insmootd through the containes. The former of these is regarded as pointing to the presence of a benzene modeus in strychume, which nucleau in breme as dimethocythated. 4H.O)¹ When moderately heated the crystals melt and loss their water. According to Guy, brunne melts at 115°, and sublimes at 204° C, the sublimate being usually amorphous According to Glaus and Rohre, after drying at 150°, brunne melts at 178°.

Bruene is more soluble than stychnine in water, dissolving in 1050 parts of cold, and less than half that proportion of boiling water. In alcohol it dissolves very readily, a fact which is employed to separate it from strychnine. Bruene dissolves in 4 parts of chloroform, in 440 of ether, in 60 of benzene, and in 120 of petroleum spirit. It is insoluble in fixed caustic alkaliss, and only sparingly in access of ammonia.

Brunno is a weaker base than strychnine, but is not extended from accludated solutions by immuneable solvents. It resembles strychnine closely in its general characters, but is less poisonous, from 7 to 10 parts of brueine having the same physiological effect at 1 part of strychnine³ II is excerted far more rapidly than strychnine, so that when given by the stomach it produces hitde effect, though it is fatal when nujected hypodermically (T. La ut do r Brun to n, Josu Olean, Soc., xivi 143)³ Like strychnine, it is not acted on reality by cold sulphurus each, or by caustic alkales. It dissolves without decomposition in strong hydrochloric acid, and forms readily crystalliasible and soluble saits

normal results of resolution at a clouds as the contract of bruche, of present proximates, but again dissolves, forming a rod solution from which d in this of write $(R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm out}) = (R_{\rm out}, R_{\rm o$

ANALATICAL CHARACTERS OF BRUCINE.

1 Brucine is precipitated in a free state on adding an alkali to the solution of one of its salts, and may then be taken up by

From analyses of their platinous compounds, Koefoed is of opinion that commercial brucine contains two homologous alkaloids

² According to Talk, the physiological activity of strychnine is 88½ times greater than that of bucune

* T. J Mays (Tow Physiol, vin 391) finds that, when frogs are poisoned, brucino primarily affects the posterior and strohima the anterior extremities; convulsions occur very early and invariably before death in strychime poisoning, and very late or frequently not at all in biguine possoning.

agitating the alkalnae liquid with ether-chloroform in the same way as strychnine (see page 364)

- 2 Brueine forms a soluble chromate, a fact which is occasionally used to separate it from strychinine A better separation is effected by crystallising the free alkaloids from hot alcohol, or by converting them into ferrocyanides (page 366)
- When treated with concentrated sulphuic acid¹ and an oxidising agent, brueine does not give the coloured products so characteristic of strychnine
- 4 The most satisfactory reaction of bracure as that with nitice and on adding a drop or two of cold nitice and of 142 sp gr to an other-chloroform residue, or other solid produced, which on heating changes to yallowsh red, and finally to yallow? If the mixture be now cooled and treated very cautiously with stannous chlorads (or other reducing agent, such as sodium thosulphate), a purple coloration is produced, which is destroyed by excess of either nitro and or the time sile.

The sed coloration of bruenne by intric and may likewise be developed by insolving the alkaloud in strong sulphiume and in a test-tube, and allowing nitric send to run on to the surface of the heaver liquid. A red zone, beaming to jellow, will be produced at the junction of the two liquids. If cold mitric send be added to be junction of the two liquids. If cold mitric send be added to be their lirgidy 'dilititled with' 'water, a body colled 'k ak to tel in, Cattagn(NO₂)MyO₂, separate in rejlow flocks: The filtered liquid, after neutahisation by ammonia, gives a precapitate of fedition collection of the precapitated laketelium may be dissolved in dilute hydrochloric acid, and crystallised therefrom in onage-red or yellow scales.

The production of a red colour with nitre and, accompanied by a formation of oxile acid and yellow scales or crystals, insoluble in water but soluble in dilute acids, constitutes a combined reaction which is peculiar to brucine

- 5 Bincine dissolves in chlorine-water with red colour. On evaporation, dichlorobrucine, $C_{23}H_{24}Cl_2N_2O_4$, remains as a reddish brown, amorphous mass
 - 6 Potassium bichromate throws down from solutions of brucine
- According to some observers, strong sulphure and imparts to brucine a rose colour, which changes first to yellow and then to yellowish grown
- Strychnine, on the contrary, gives no coloration with cold nitric acid, but developes a yellow colour on warming
- The orange colour produced by adding intric acid to morphia remains unchanged on addition of stannous chloride

salts, even when very distri, a yellow precupates of bruches chromats, mesoluble in meter acad, but salable with deep set of comin strong intre and. The microscopic appearance of bruches chromato as characterizates, and, togother with its behaviour with nitre each, distinguishes the precupitate from all others produced by the reasent

7 The microscopic appearances of the precipitates produced in brucine solutions by platinic chloride and potassium ferricyanide are also highly peculiar

8. Potassium ferrocyanide only slowly precipitates acidulated brucine solutions, and affolds the best means of quantitatively separating brucine from strychnine (page 366)

Nux Vomica, 1 Poison-nuts. Quaker Buttons

The seeds of Strychnos mux vomica are known by the above names. Their appearance is highly characteristic. They have no odour, but taste intensely bitter ²

If powdered nux vomes seeds be moistened with water and examined with a low power, the characteristic fibrous hims can be readily recognised. They sequires a yallow colory on adding todased podasesum oolde, while he rest of the powder becomes burn Touched with strong intense and, the powder acquises an orange-red colory, rawhally uesterwed on adding stanous chiracte

Nux vomice contains, in addition to the usual plant-constituents, the posonous alkalaris strychnine and brucine, a glucoside called loganin, and a peculiar and named strychnic origasuric acid

STRUMING OF IGASURIO AGID appears to be a variety of famini, It was obtained by Hohn (1873) as an amorphous yellowishwhite mass of strongly and and somewhat astringent taste. It gives a dark green coloration with ferric chloride, a white prequiptate with lead acstate, and mould reduces amouncements of silver.

¹ French: Now vomuoues German, Krähenaugen, Brechnuss,

² The powder of nur venmes has a grayah-buff colour, and, in the experence of the author, has been sold by a registered druggest in mistake for julap. Death has been caused by the sale of nux vonice for hydroice lowels, which, by attificial light, is of somewhat similar appearance (Pharm. Jour., 13), vo. 401.

2 It a microscopus section of nux vennes be teased with potolemn spirit to remove the fat, the parts containing brunes will then assume a bright red colour on being moutened with a maxter of seleme and intro seeds. To detect strychima, the section is treated in sencession with patrolemn spirit and absolute acchole to remove segar and bremosy, and then tested with a solution of cerum sulphake in sulphurne said (O. Li in d. Chem Centr., 1884, page 489).

Loganin axish in mur vomice seeds, but more largely (4 to 5 per cent) in the pulp in which they he embedded in the furth Dunstan and Short ($Phanm\ Josu$, [3], xiv 1025) obtained loganin in prasmatic crystals by cooling the liquid obtained by exhausing this substance with cholorform-alcohol (4 1). After re-crystellization from alcohol, the crystals contained $C_{\rm SH_2}C_{\rm QL}$ on supprised formula identical with that of ar but in, from which, however, loganin is distinguished by its much higher moltang-point (above 200° C), and by not verifying common with dilute subluture above

Logann is seadily soluble in water and alcohol, but less so in either, chloroform and benzene. It developes no colour with interior and on other excitising agents, and the aqueous solution is not affected by solutions of lead, iron or silver, and does not reduce Fehling's solution. When gently warmed with stong sulphure and, loganin gives a fine red colour, changing to purple on standing. When bolled with ditute sulphure and, it yields a colouring gluous and logan stin, which latter body behaves with

solvents and reagents very similarly to loganin steeff. For the cases of nur vonues, D u is tan and S h or t (Pharm Jour, [3], xin. 665, 1056) tecommend that 5 grammes of the finely-divided seed (previously dised at 100° O) should be exhausted in a Soxhlet tube or its equivalent, for one or two hous, with a mixture of 40 or of chloroform and 10 of stooloh! The solution is againsted with 25 or of ditule subplume and (5 per cont.), and the chloroform separated and again agitated with 10 or of ditule and T has separated and solutions is affixed, if necessary, rendered alkalime with summonia, and chaken twos with chloroform, using 15 or each time. The chloroforms oblition is separated, filtered, exponented, and the residue direct at 100° for about an hour, or till constant in weight. The following results were obtained.

Description of	Sam	lo		Date of Collection	Total Percentage of Alkaloids
Bombay, fine, Bombay, ordinary, Bombay, Cochin, Cochin, Madias, Madias, Avetage,	:	:	:	1877 1877 1883 1887 1888 1877 1888	3 46 3 14 3 90 3 04 8 60 2 74 3 15

² This mixture is described by the authors as one containing 25 per cent of alcohol

The alkaloid in powdered commercial nux vomica ranged from 2.56 to 3.57 per cent 1

Ether-chloroform may be advantageously substituted for unixed choroform un the foregoing monosa and the alkalonds may be conveniently titrated with a standard memeral and and methyl-causing instead of being weighted. One co of decunormal sead neutralises 0.0364 gramme of a mixture of braces and stretchine in molecular proportions of 333 4341, When deared, the strychinne and buceine may be separately determined as described on new 357

Extraor or Nux Vontos, BP, is disected to be prepared by exhansing the duest and powhered seeds with somewhat diluted spirit (4 1), and evaporating the filtered higher Foundry the extract varied considerably in strength, invelve specimens of the commencial article examined by Du nation and Shori in 1844 (Phaim John, [3], xir 443) containing proportions of 1644 alkaloids ranging from 10-32 to 17-54 per cent, while the ratio of strychime to bracine varied from 1 1 up to 1 17-9. On the other hand, the proportion of water only varied between 13-6 and 19-7 per cents².

In the Pharmacoperus of 1885, the exteact is directed to be standardsed so as to centuan 15 per cent of total alkiloids. For its assay, 10 grams of the extanct are directed to be dissolved in assay, 10 grams of the extanct are directed to be dissolved in 3 grams of sodium carbonate m $\frac{1}{2}$ or of water, heating gently if necessay, and a solution of 30 grams of sodium carbonate m $\frac{1}{2}$ or of vater added. This solution is then agatated with $\frac{1}{2}$ or of chiloroforms, which extracts the alkalouls. This iteatiment should be repeated (Pharm Jow. § 3), xxx 625), After which the chloroform is shaken with dilute said, and this solution extracted with chloroform and ammonia in the manne already described (page 385)

As strychnine is greatly more active than blucine, and the relative proportions of the two alkaloids in the extract are by no means constant, it is questionable whether it would not be pre-

¹ Powdered mu vonuces has frequently been dued at a temperature above 100°, in which case the oblisoform alsohol centre often continus colouringmatics which illumetally contaminates the alkaloid. In such cases the brown, colour may be knowed by againing the oblisoform-alcohol solution with an aqueous solution of crystallised sodium carbonate (5 per cont.) before treating it with dultae on.

² Bockuits (*Pharm Jour*, [8], xx 241) found in five samples of nux vormes extract the ratio of strychnino to bracine varied only between 48 57 and 54 46

8 The absence of relation between the total extractive matter and the alkaloids of nuv voninear renders the official method of standardising the extractiver unsatisfactor v

ferable to ascertain the proportion of actual strychnine rather than that of the total alkaloids.

G F. Schacht (Phan m Jour, [3], xw 85), recommends for the rapid assay of nux rounce extract that I gamme be disolved in 30 ec of water, the solution andulated with 1 cc of hydrochlone send, warmed gently for half an hene, and allowed to cool. It is then filtered and made up to 100 cc. Ten cc of this solution is then tituated with \(\frac{1}{160}\) Mayer's solution (page 139), such cc of which represents 000184 gamme of mixed siychinus and brunne \(^1\) The results by this process are stated to agree closely with those obtained by the gravimente method.

Nux vomica extract contains from 9 to about 20 per cent of water, and some specimens lose and others gain weight on exposure

TINGTURE OF NUX VOLUGA, BP, is directed to be prepared by dissolving the extract in slightly diluted sparts, as a to contain 1 grain of total alkalouls in each fluid ounce, equivalent to 9.29 grain per 100 measures. Dun a ten and S hort (Year-Dool. Plan as, 1883, p. 476) found the specific gravity of twelve commenced lunctures, obtained from the principal London manufactures, to vary from 8.377 to 8593, the proportion of strychnime from 0.46 to 0.131, with an average of 0.080 per cent, the human from 0.45 to 0.239, averaging 0.130 per cent, and the total alkaloul from 0.124 to 0.360, with an average of 0.218 per cent Before 1885, when the tinuture was directed to be prepared from a duly standardised extract, its strength was very variable? The tinuture of nux vonus may be assayed by evaporating the spirit from 60 cc, treating the readuce with dulute sulphurue and and chloroform, separating the read, and extracting the alkalouds by ammonst and chloroform.

Alkaloids of Curare.

The Indian arrow-poison known as curare, curari, wourali, weorara, or usali is a possonous extract prepared

 1 The solution is prepared with 1 355 gramme of mercuric chloride and 4 98 of potassium reduce in the little

The pupuation of the tracture of new roomes from a standardand extended has agreement finded to secone undermuty in the compression, for of tway four amplies of the commercial interiors purchased by N IM Na **11 n m 1889.88, mm routiles after the publication of the new edition of the Flower-most elevent showed by them pale yellow colour that they had been prepared by the old process from the needs, and contained from 0.110 to 0.885 par cent of total alkaloids, while the processings of total alkaloids in the thintoon samples which by their light brown colour showed they have been prepared from the orticast, ranged from 0.196 to 0.313 pcr cent (Yeon-Dook Phorm , 1886, page 507)

The intensely active arrow-poison used by the pigmies met with by

388 CURARE

from the bark of Strychnes tax/fors, a native of Guina, together with other vegetable extracts. It coems no commerce as black, shaming, buttle, resmond mass, of an intersely bitter taste. About 83 per cent is soluble in water, and 79 in distuled spinit. A maxture of glycerm and dinited spinit disturbed of sylverm and dinited spinit disturbed of sylverm and dinited spinit disturbed of a fixed Curare, as imposted, varies much in strength, and often containes calcium eubonasis and phospitale. It is exceedingly possioners, and should be handled with the utenot care. Curare should never be allowed to come in contact with a cut or sentich, and, unleed, hould never be touched with the naked fingers, or powdered or monumistical in the chry state.

"Much doubt exists as to the tran nature of woorars. According to Waterton it is prepared from several different plants, two species of possonous ants, and the fings of cortain snakes, while Schom burght states that it consists of vegetable institle alone, and chiefly of the bank of Skyalone toutprin. That there are at least several varieties of this substance current among the different tubes of Indiana seems to be fully established. , and it is even probable that each trube has its own method of preparing the poson" IT G W orm It e. Naco-Ofemistry of Pausons'

Cumune excreases both a puralysing and totanium gration, but it special to our tit chief poissions properties to its action on it is necros of motion, which it paralyses, so that an animal under its influence these of sufficient of from puralyses of the muscles of the chest. Hence its physiological effects closely issemble those produced by methyl-staychima. According to J Tillie, when the difficulties besetting the examination of the storion of curare on the spanie could are avoided, curare produces totains just less trychima. Carare appears not to set as a posson when taken into the stomach, but when employed as a hypodorme injection of 15 grain has been found fatal to a rabbit, and 004 grain to a frog If, after administration of curren, life be mantened by arthorize respiration, symptoms of diabetes mellitus are observed, and the union is found to contain energy.

Notifies strychmue nor bruene has been detected in curare, and that the paralysing effects of the preparation are not due to methyl-stychmue is apparently proved by the superior texicity of the vegetable extract J Tillio (Jour Anat and Physiol, 1990) attributes both the paralysing and tetanising action of curare to

H M Stanley in Contial Afron is compounded from five plants. Its toxic action is believed by E M Holmes to be due to orythrophicoine and strychnine (Pharm. Jour., [3], xxx 917)

¹ See a valuable paper by J Moss, Pharm. Jour, [3], vin 121.

CURARINE 389

cutarine, but it seems not improbable that the preparation contains at least two active alkaloids, one having a paralysing and the other a tetanising action (as is the case with Calabar bean).

Curare has been proposed as a remedy for hydrophobia and as an autidote to poisoning by strychnine

CULALINE is the name given to the physiologically active base of curase, and the imploable formula C₃H₂H₃ has been sarrhed to it. Curatine has been variously described by different observes and it appears exchain that the products have been of very discordant chanacters. Curarine is described (1865) by Preyer (Ohem. News, xii. 10) as cyt-talliang in very hygroscopic four-sided prisms, axing a bluter taste, freely soluble in water and alcohol, only sightly so in chloroform and amyle alcohol, and insoluble in ether, burners, turner tune and carbon disablished.

The zyueous and alcoholic solutions of curarum have a bitter taste and family alkaline seaton. The base is said to fore crystallisable salts with hydrochloric, nitric, and acetic andisfric commercial curarum prepared by Menck, according to Dehical inchhol, is described as a yellowsh brown, amorphious powder of nitensely bitter tasks, easily soluble in water and alcohol, but insoluble in ether I shows no perceptible alkaline reaction, and forms no true salts, but on evaporating an aqueous solution in diltte seid to a syrup, secular crystals of an inactive decomposition-product are formed, whereas the lethal dose for gumea-page of curarum etself is stated at 0 00035 gramme per kilogramme of weight. Concentrated sulphure and dissolves Menck's curarus with crimson colou, changed to blush by podassium bachromate

With strong nitric acid Preyer found curarine to give a purple coloration, and with concentrated sulphunic acid a magnificent and lasting blue colour C Bernard found the colour with sulphuric acid to be a carmine-red.

If a filtered and highly concentated solution of cunarue be mixed with dutue giverin, and a saturated solution of potessium inchromate added, amorphous curarine chromate in precipitated. Even after solution in boiling water it is again deposited in amorphous state, a fact which distinguishes it from strychume

¹ That the tetanising action of curse is due to the species of Shyahine employed for its preparation, and not to generoteen or other principles district from the various plants sometimes used in conjunction with it, as proved by the fact that a genuine specimen of the bank of Stephone tempers produced the same symptoms (I Tillie, Jose Anat and Physici, XIV 42, see also Nikolski and Dografs, Per-Book Pharm, 1920, page 188)

² According to Sachs (Annales, exc. 254), Preyer's crystalline curarina sulphate consisted of impure calcium phosphate (1) with mechanically adhering curarine chromate, which forms well-defined crystals Curarine chromato is more soluble in water than is strychnine chromate, and is nover perfectly precipitated even by addition of glycerin or alcohol

If the pocupitate of curainne chromate be kept for some atmost decomposes, but if totald without delay with concentrated aulphures acad it developes a magnificent blue colour, which is often wolet in the presence of impurities (P el 1-can observed a brilliant set obloates) in The reaction simulates that obtained in a similar manner with strychnine, but curamic can be separated from strychnine by rendering the cold solution alkaline with ammonia, and then filtering Strychnine will be found in the prequiptate, whilst the curame will remain in the liquid, owing to its solubility in water. The filtrate may be agitated with oblicoform or benzene to iemove any time of strychnine, the aqueous liquid concentrated and the curains converted into chromate and tested futther, as already described.

Curaine is very unstable, and hence its solution should be subjected to as little manipulation as possible

CURING exists, according to Bohm (Ber., xx 143), together with curarine in many specimens of commercial curare. Curine is said to exist in the aqueous extract of curars, though in some cases dilute sulphune acid is requisite for its complete solution. Upon rendering the liquid slightly alkaline with ammonia, a dirty green precipitate of impure curine is found, which, by successive purifications with ether, alcohol, and again with ether, may be obtained in a micro-crystalline condition. Curine is described as melting at 160° to a clear liquid, and being slightly soluble in cold water, freely in alcohol, chloroform, and dilute seids, but less readily in other. The most characteristic reaction of curine is the formation of a voluminous white precipitate with metaphospholic scid Curine itself is stated by Bohm to be physiologically inactive (in doses of 5 to 10 milligrammes), but by treating it with methyl todado he obtained the hydrodade of a new base which possessed an intense curaio action, I milligrammo killing a guinea-Dig (weighing 1600 giammics) in one hour J Tillie (Journ. Anat and Physiol, xxv. 42) states that curine has no apparent action on motor nerves, but when hypodermically injected acts on the hearts of both frogs and labbits as a paralysant similar to veratime or digitalis. As curine is liable to be present in curere in very variable proportions, its possible presence in commercial curarine must not be overlooked

CINCHONA ALKALOIDS.1

The virious species of the family of plants known as the Conchonacea yield an extraordinary number of closely analogous alkaloids. These bases exist chiefly, though not wholly, in the bank of the trees, and are remarkable for their valuable febrifuse properties.

The constitution of the cinchona bases is at piesent very imperfectly understood — Quimamine and cuprenne are known to contain hydroxyl-groups, and quimne and cinchonine and their isomors have been proved to be derivatives of quinoline. An abstract of the existing knowledge of the subset is given on ragic 189.

Any axis-factory classification of the emchois bases in the present impurites state of our knowledge of their constitution, and in some cases even of their empurical formula, is manufactly impossible Loneauenn is common, and on slight provocation quiume and some others suffer polymenistics, with or without leaning the elements of water, forming memolipous "npo" of antivito-based

Perhaps the most suggestive method of classifying the emchona bases and their allies is to airunge them according to the number of atoms of oxygen in the molecule, and subdivide these classes according to other analogies

The following (pages \$92, \$93) as tabular last of the alktolucle bulberto selated from the various species of enchona and alled bulbs. It contains the names of all the natural emchans bases, the extrience of which as chemical individuals has been fauly well established up to the present time; but it must not be supposed to include all that actually exist

As as evident from the table, somenium is very common among the cambnen bases. Thus the two best-known bases are quisars and emchonime. Isomerides of these bases coanst with them in the balk, and are called respectively quinatine and candentitine. It is probable, however, that the base neadly teamed enchouldine presents the closest parallelism with quimne, and that emchomine is the analogue of quiumline.

The four bases above mentioned are the chief cystallivable alkaloids of embona barks, but there exist with them, or are formed in the process of manufacture, certain amouphous isomerable called respectively quincine and enchloratem. It is doubtful how far these bares pre-exist in the bank, the natural amouphous alkaloids being probably the anhytro-derivatives dispusions and

 $^{^1}$ The author 14 indebted to Di B H Paul and Mr A, J Cownley for the perusal and correction of this section

TABLE OF CINCHONA BASES

Alkalord	Formula.	Chast Source	Melting Point,	g Optical Retation, Sp 1	Other Characters.
I CINCHONINE CLASS— Parlome, Cinchotme,	CleHusNgO	0 lutes and C succerubra from Darpoling Crude circleonie sulparte	81 12	• +	Pale yellow, amorphous, bitter powden, Sparingly soluble maste Slonder prisms and soules.
Hydroenchonne, Eydroenchonne, or	OgH34N30	Compute Particana. Compres Mother hunors of homoun	5 58 5 58	+1311	Yery posonous floragonal prisms. In fanlandoun reschon Tage 488. Yellow, amorphora powier Plates or flat needles. Not fluorescent.
Carchonae, Carchonae, Carchondine, Homosmebondae,	Cashen NgO	Chondine Various species of Chackong Almost always passent. Repectally O rubra With cinchonique.	200	+226 6 -70	Rhombic prisms. Page 431. Prisms Page 423 Prism or plotes Not fluorescent No
Chrohonteine, Paytine, Paytanine,	O _E H ₂₄ N ₃ O	By heating cinchonna, From winto bark of Rayta.	88.	1881	thalleadum raction. Amorphous, Crystallisable salts. P 485 Crystallises with I agus in fine prisms Amorphous. Amorphous salts.
II. QUINAMINE CLASS— Quinamine, Conquinamine, Javazine, Chyrene,	C ₁₉ H ₅₄ N ₃ O ₃ C ₁₉ H ₂₆ N ₃ O ₂	(G successfort from Brit. Indus and Java. and Java. (C officiation Live. C celescys from Live. C capture, or Remajos pademortales.	25 E 85	+101 6 +201 6 -175-3	Long prama. Page 627 Long channa, trieline presse. Page 427 o Ricombe Diskes. Soluton in H ₂ SO ₄ intensely pollor. d Grouped primma. Page 438.

TABLE OF CINCHONA BASES—continued

Alkaloid	Formula,	Chief Source,	Meltang Optical Point, Rolation Sp.1	Optical Rotation, Sp.1	Other Characters
III QUININE CLASS-					
Hydrogumidme,	Cau Hai N2O2	In mother-liquors from qui-		-142 2	Bitter meedles, Fluorescent, That-
Qumme, Quandme.		Co-frame surprise		+ ;	or tables Fluorescent reaction.
Quinome,	forestana. ∫	By heating quame sulphate.	:43	1588	Phys 397 Phys 425
Charamine.	_		_	!	Americans of oil. Non-fluorescent P 434
Concharramine, Chairamidhe, Concharramidhe	CmEnN304			+100 +68+	a Fredles or prisms containing 1 agua.
Concuscomne,	Call w N.O.	Romma Purdicana (Cusco	EEF	282	a Amorphous powder a Orstallactife
Спасовине,		or came Onpres Bark) .	_	23	a. Non-bitter, shining prisms Acetate
Cuscondine,	CoHasNo.		011	-543	Monochant plates with 2 agus Not
Cuscamine,		C Pelleturana	(213		Yellow, amorphous
V AKHTDRO-BASES- Dromchomome,	CoH.N.O.				· ·
Dagameine.	Col. a. o.	c. rosmenta and O sucer-ubra	9	\$	Yellowish, amorphous No then amorphous
	SOT STATES	C resulents and "quinoidine."		+	# 4
			_	_	

¹ The values for specific relatory power refer to solutions of the free alkaloids in nearly absolute alcohol

dicinchonicine, and distinct from the amorphous products formed from the crystallisable bases by the action of heat or acids

In addition to these isomers and anhydro-derivatives of the cinchona bases, there exist various homologues and isologues of them. Quinine itself is probably a methyl-cupieine and a methyl-cupieine and a

Catam of the cinchous bases (eg, cinpreim) exhibit a remarkable tendency to form stable crystalline compounds with other of the bases. It is probable that the existence of these remarkable compounds, having different physical properties in the form of salts as well as in the fire state, has led to the isolation and description of various bases which will hereafter be found to be compounds.

The less impotant emchona bases have no recognised position in commence on mediune, but they are label to be present to a greater or less extent in specimens of commencial citaloris called by the better-known names Commencial citaloris called the retain traces of cinclonine, quinding adjunctly contains notable projections of emchonaline Hydrocinchoudine is sometimes present in commencial or in ch on rid in a while qui in it in contains hydroquination and hydroquination. Quinamine and conquinamence are probably not unfrequently present in commercial cancions alkadoris and configurations.

General Properties of Cinchona Bases.

The circhona alkalouds all have well-defined basic characters, some of them being sufficiently powerful to displace aumionia from its compounds. Their salts are usually crystallisable

In the free state, the emchema alkaloak are colour loss of familyyellow solisk, often readily fivules, but not volatile without documposition. They have generally but little solubility in water, but dissolve more readily in alcohol, and generally write great facility in ethes and chloroform. Such as are soluble in the last two liquids are nonwest from their ammonancal solutions by agitation with ether or oblivoform, but in no case will either or oblivoform emove them from an aqueous solution acculated with sulphume or hydrochlorio acad. On the other hand, the anhydrous sulphates of many of the cambons alkalouls are soluble in chloroform, and still more readily in a mixture of chloroform and absolute alcohol. This fact is sometimes uthissed for detecting adultivations (p. 417)

The solutions of some of the cinchona alkaloids in excess of dilute sulphune acid exhibit a strong blue fluorescence, which is visible even in very dilute liquids. This fluorescence is destroyed by adding an excess of chloride of sodium or other haloid satit. The solutions of the cunchona alkaloude exact a well-marked rotatory action on polarised light, the colution being in some sessinght, and in others left-hended. The specific rotation is affacted in a remarkable manner by the solvent employed and by the proportion of free and present, which circumstances greatly reduce the practical value of the optical activity for the identification and quantitative determination of the unmixed lakalouds.

On adding a fixed alkalı, alkalıne carbonate or ammona to the solution of a salt of one of the cunciona bease, the syrengty soluble alkaloid is usually expanded in a free state, and is in some cases soluble in an excess of the precupitant. On agatantig the alkaline liquid with chloroform, the precupitant of angatantig the assolved, and may be recovered in a fire state by separating the chloroform, and oraponating it to thyness at a steam-heat By adding more schloroform to the aqueous lequid, and inspectating the agitation, the complete extraction of the alkaloid may be ensured, and the process made quantitative (see page 419). Eitler may be substituted for chloroform in the case of quinno and other alkaloids reachly dissolved by the discount of the case of the complete cases.

The emchana bases are tertiany amines; for when treated with an alkyl added they form additive-compounds which are conveited by treatment with oxide of silver into powerful soluble bases analogous to the telethyl-aminonium bydroxide (page 19)

Maxy of the enction althous form two sense of sails, neutral (improperly called "base"), and acid sails. The neutral sulphates of the emclions alkaloads have, when subpriore, the general formula R,H,SO. They have a neutral reaction to littens and methylorange, and are generally very sparingly solution in water, but corresponding acid or breatlyhates (BH,SO) are generally readily soluble. In some cases still mose and sulphates are known

The sulphates of many of the anchona bases possess the property of combining with solane, the compounds produced being m some cases of a very complex character. Certain of these "1 odosulphates," of which the gumma compound or heropathits is the type, possess the remarkable optical properties of the tourmains (see mage 103)

When a salt of one of the natural cunchona bases is heated for a pulonged person to a high temperature, the skiladou undergoes a curcous change. It becomes incapable of anystallising, a property sometimes extonding to its salts. The change occurs most readily by exposing the need sulphate of the skiladoud to a temperature of 100° till analysidous, and then innecessing the heat for some time

³ This is not the case with cuprome and some other alkaloids, which form definite compounds with the fixed alkahes in the same manner as morphine

to about 130° C No means are at present known by which the modified alkaloid can be restored to its original crystallisable condition

When the cunchons bases are heated with stong hydrochlore and (sp gr. 1 125) to 150° for six to ten hours, they are converted into apo- or anhydro-derivatives of bease character, the change in the case of quinne and quinidine being attended with ovolution of methyd chorde (H os so, Annal, cov. 314)

When the sulphates of quame, canchonine, and cutebourline are dissolved in concentrated sulphune scale at the endmary temperature, they are converted into "iso-bases" (Hesse, Annal, cextin 131), which differ in several respects from the piracutalisations Hydroquamine, part and hydrocunchonidate are converted by the same treatment into the corresponding sul-bh nine acids, which are commounts of distinct base characteristics.

With platimic chlorids, the hydrochlorides of the cinchona bease form chloropplatinates of the genoal formula BH₂FCI₀, but many of them also form salts containing LTLFCI₀. Salts of this constitution are produced on adding sodio-platinic chlorid to neutral solutions of quinne, quinchine, cinchonidine, and homochonidine (Hesse, Amad, everu 929) The auro-chlorides of the cunchonides proposition with soparation of finely-davided metallic gold speedy decomposition with soparation of finely-davided metallic gold

Certain of the einchone bases give a deep green coloration or precipitate when their solutions are treated with chlorine or bromine wator, and ammonia subsequently added This reaction is known as the "thalleloquin test" (see also page 401)

Most of the cinchona bases are very completely precipitated by tannic and piene acults, potassio-mercuric rodule, and certain other reagents. These reactions are sometimes used for their dotection and separation

On treatment in solution with bromme-water in slight excess, the cuncions bases are converted into brom or -derivatives. The number of atoms of bromme taken up varies with the constitution of the slikeland According to TF aww soit (Pharm Jour, [3], xx. 915), quimine, quindine, and eupreime react with approximately Bi₀, hydroquinine with Bi₂, and cuchonine, cuncloindine, and "anouphous quimine" with Bi₃ on basing the eight occlude, and the sum of the same properties of the same properties. The same properties of the same properties. They can be drugted at 100° without change, are readily soluble in

dilute acids, and are thrown down as reamous precipitates by alkales. On treatment with alcoholic potast they are hydrolysed into acctic acid and the original bases. The acetyl-derivatives of quinne and quinding give the thalleroquin reaction.

The more important properties of the leading cinchona alkaloids may be summarised as follows .---

(Hydrated crystals are formed by Quinine, Quinidine, Paytine, Cupreme,

Anhydrous crystals are formed by Chichonno, Chichondhio, Quinamine.
No or ystals are formed by Pancine, Quincine, Diquincine, Dicinchonicine
(Readrly soluble in Ether — Quinine, Quinamine, Paytine, Quinome, Java

B Sparingly soluble in Ether —Cinchondine, Quinidine, Cupreine

Dezir e-rotatory solutions in alcohol are formed by Cinchonine, Cinchonamire, Quinamire, Quindine, Chananire, Quineme, Diquinione, Chananire, Quineme, Diquinione, Chananire, Quineme, Diquinione, Chananire, Quineme, Diquinione, Chananire, Chan

Lawa-rotatory solutions in alcohol are formed by Cinchemdine, Hydroemelousdine, Homoenchondine, Paytine, Cupusus, Quinne, Hydroquinine, Casconne, Anguso

(Fluorescent solutions in dilute sulphune acid are formed by Quinnie, Ounning, Hydronium, Hydronium, Diquinicino

D No fluorescence is exhibited by solutions of Cinchennia, Cinchendine, Hydrocinchenidine, Homocinchenidine, Quinamine, Quinamine, Quincine, Dicin chonicine, Cusconine, Cupieine

Thallesquin is formed by Quinne, Quinidine, Quinione, Diquinione, Hydrograms, Hydrogramidine, Capticine

El Thallesoguin is not formed by Apoquimidine, Cinchondine, Cinchondine, Homocinehonidine, Hydrocanehonidine, Cinchonicine, Quinamine, Cinchonamine

Quinine. Quinta

determination in the same.1

Quinne is the most important of the cinchons bases, and appears to possess the most important of the cinchons bases, and appears to possess the most powerfully febringe properties. Its mole of preparation from the bark is based on the same principles as its

The final-y-powdered bask is ground to a time pasts with lime, caustrosold, or section excitonate, and exhaused with warm possible oil of separates, when it is in nof and shaken with sulpharmon cold, this solution is boiled, and withst boiling is nestinated with solution can beaute and solution to cold (write with to being in sectional solution with consideration concentrations, and committee crystallizes ont on cooling, which cardiomains calciume, and committee cardiomate results in solutions are substant in sulphates. The quantum chances plate is purified by recrystallization after iterations twith annual chances of the contract of the contract of the contract in the contract includes a solution and the contract of the contract includes a solution and the contract of the contract includes a solution and the contract of the contract includes a solution and the contract of the contract in the contract solution and the contract of the contract o

The chemical constitution of quinnic is not thoroughly understood, but such knowledge as exists is optionised on page 168. The complete synthesis of the alkaloid has not hitherto been effected, but cupretine has been appearently converted into quinniby the introduction of a undhylytgoup! Two distinct bodies isomeric with quinnic have been synthetically prepared (page 169).

Free quinine usually appears as an amorphous or resinous mass In commerce the tree alkaloud is usually met with as a coarse powder, having a brownish yellow tuit owing to a trace of colouring-matter. It may also be obtained as a fine white powder

From alcohol and some other solvents quinne may be obtained in crystals, but on the ovaporation of its ethereal solution it separates as a goldations or resented meas, which is never crystalline. This behaviour is unportant, as most other eincliona bases are crystalline ether-iesaform.

As obtained by the precupitation of one of its salts by an alkalt, quantile forms a bulky, white precipitate, which congruences have not a resemble and the procipitate at first formed at the ordinary temperature is amorphous and anhydrous, but it soon takes up water and becomes crystalline. It then contains 3 aqua. If the ammonia be added in large evcess, and the solution is not to concentrated, the trib yld rate is obtained in small needles.

and extracted with weak alcohol Of the three bases precipitated by the akaki, quindine and cuchomaine are dissolved by the spirit, whilst cinchomine is left behind, the two former can then be separated by means of their neutral tattrates, that of quindine being considerably the more spirible

Grimanx and Ainold, Compt. Road, exil 774 When a solution of

cuprenes to methyl alcohol is bothed for several home, a dan 774. When a solution of cases, with the theoretical quantity of schulm and occess of mothyl todale, a maximo of new adomethyl stew was obtained, having all the chanctons consumerable resulting from the similar treatment of quantum. By substituting mostly claim for the todde, and operating in a sended tube at 100%, a base section, the sulphate of which had all the chemical and physical characters of quantum sulphate, the following teaction having probably occurred;—

 $C_{19}N_{21}N_2O$ ONa + $CH_3Cl = NaCl + C_{19}H_{,11}N_2O$ OCH₃. Sodium compound Quinte and the same compound can be obtained from an ethereal solution below 10° But the resmoid mass left on the spontaneous evaporation of a solution of cumine in ether usually contains water in proportion corresponding to a monohydiate, and when the crystallised trihydrate is exposed in an exsicentor over sulphuric acid, it offloicsees and loses its water more or less perfectly 20° C, over strong sulphune acid, the trihydrate soon loses the whole of its water, but over equal measures of strong sulphuric acid and water a monohydrate results. At 15° C, in the open air, the triliydrate is unaltered, but at 20° C it effloresees and loses I aqua, the residue having the composition of a dihydrate Commercial quinine contains from 8 to 11 per cent of water, and lience is approximately a dihydrate. The precipitate produced by ammonia at a low temperature in concentrated solutions of quinine sulphate is also usually a dihydrate. Hydrates of quinine containing 8 and 9 aqua have also been described. When the tuhydrate is exposed to a temperature of 40° for a short time, and then to 60°, the whole of the water is driven off, and this change occurs rapidly at 100°. Resmond gumme loses its water with some difficulty at 100° unless previously powdered, but at 120° becomes anhydrous very rapidly (see Pharm. Jour, [3], xvi 385. 897, 937)

Auhydrous quinine, obtained by drying the triliydiate over sulphure acid and heating to 115°-120°, melts at 171 2°-172°, and that prepared by heating the benzene compound to 120° at 171 6°-172° 1

Quuune is odouless When in solution or finely-divided it has an intense and purely bitter taste. It has valuable fabrifuge properties, and is possenous to frogs and other of the borramains. It has decided antisophic properties, retarding or arresting the alcoholie, lactic, butyric, amygadious, and salicylous fermentiations, but not the dige-stree action of pegsin.

Quinne is very spannigly soluble in wale, according to J Regnauld the solublity at 15° C being I pat in 2024 According to Sestin, however, the solublity of the unlyvious alkaloid in water is 1 in 1667 at 20° and I in 902 at 100° C, the tribydrate requiring 1428 and 773 parts of water at the same temperatures.

In dilute solutions of the fixed alkalies quinme is not more

According to Hesse (Annal, colvum 133) on prolonged heating of a solinoid quantie in alcohol to 30° the alkaloid is converted into an isomeride for which he proposes the nametable name of he me quit in no. This melts at 174 4"-175°, and is reconverted into quinne by prolonged heating with dilute subhure scale.

soluble than in pure water, but ammonia exercises considerably greater solvent action. Certain ammonium and calcium salts notably increase the solubility of quinine in aqueous liquids.

Quinme desolves in shout two parts of alcohol of 0.82 sp. gr. and is still more soluble in boiling alcohol Crystallised quinne is stated to sequire from 22 to 30 parts of ether for solution, but freshly-precipitated quinne dissolves in little more than its own weight of other Quinnes is also very soluble in chieroform (1.5), and dissolves readily in benzena and carbon dissiplinde its sonly spanngly soluble in petroleum sprits, even when hol.

Quinne exercises a powerful lievo-rotatory ection on polarised light, the value of S_p being, according to Hesse -145 2° -0 567 e at 15° C, for the solution of the hydrated alkaloid in 97 per cent alcohol In its salts, the optical sciurity of quinne has different values

Quinne afford no viable colour on other neactions with strong acade. By cantonasty dissolving quinne hydrate or sulphate in a mixture of equal volumes of concentrated intere and sulphures eads, amorphese d in in tro, $\Omega_{\rm sul}(R(N_0)_{\rm p}N_0)_{\rm p}$ is produced, nearly insoluble in ether and forming unerptaliastic state (E H R en n i. e, $J_{\rm sur} P_{\rm chem} S_{\rm ce}$, xxxix 489). The action of permanganate and chromic said mixture on quintine is described on page 168.

Quinine is a powerful base, its solutions having a marked alkaline reaction to himus and methyl-orange, and neutralising the strongest acids It does not redden phenolphthalein

DETECTION AND DETERMINATION OF QUININE

The detection and estimation of quinine, when it occurs unmixed with other alkaloids or organic matter, is very leadily effected, but the problem becomes more complex in the presence of other emchana bases

The following reactions are yielded by a solution of quinine in a moderate excess of dilute sulphuric acid —

1. Solutions of quinne in duties sulphure and exhibit a strong blue fluerescence. The effect is beet observed in very dutes liquids, and is intensified by addition of excess of sulphure and The hydrochlorids and other haloid compounds of quinne (including the thinsulphate and cyanogen compounds) exhibit non-scenes of solutions of the sulphate is destroyed by very small quantities of hydrochloric and or other chlorides, but can be again produced by adding excess of chitte sulphurie and Alcoholic

¹ Quinine is deposited from its solution in waim benzene in crystals containing (C_{2c}H₂₁N₂O₂), C₃H₆, 2aq (Chem. News, xlvin, 4).

solutions of quinne exhibit but little fluorescence, and solutions in the alkaloid in immiscible solvestie none at all. Under favourable conditions, the fluorescence of quinne becomes an extremely delicate test for the presence of the alkaloid. Thosescence is also produced by quandine, hydroquinne and hydroquinudine, and diquincine, but not by quinamine, sinchonine or its isomeris, onesconne, cutreine, or quinneine.

- 2 According to A Weller (Arch & Pharm, exexy 161), on adding delion-water to a strong golution of quame the solution sequere a mose or less intense red colour. Bromne-water is a strong solution of quame the solution sequeres a mose or less intense red colour. Bromne-water is a practable reagent, and on adding a few drops to a saturated solution of quame hydrochlorde a yellow prequitate is formed, which gradually disappears with formation of a nose-ord colonation, changing to cherry-red. The colour disappears after a time, but on he reproduced by adding more bomme-water, and the reaction is more deboats and picampi if the quame solution be previously gently armed. Acust and excess of bromane-water prevent the reaction, which is also produced by quandine, but not by custolous or gentleonation.
- 3. If a solution of quinne, randsaed as nearly neutral aspossible, be treated first with chloime or bromine, and then with excess of ammons, a green substance called thalloroquin is produced, which in concentrated solutions forms a prequiste, and in more dilute a deep green lequid. When corefully applied, the test, which is due to Brande, is extremely delected. Bromine is a more sensitive reagent than chlorime. The following is the best mode of applying the test—To 10 ec of the solution of quinne aid 3 ec of chlorim-water, or 55 ec of salurated bromine-water. Agitate well, and than add one drop of stong ammons solution, or sufficient to render the liquid distinctly alkaline. If the proportion of quinne exceed about 1 per 1000 of solution, a green substance is preequitately solutibe in absolute
- ³ The fluorescence of quume as best observed by holding a test tubs filled with the solution in a vertical gention before a window, whin a blussis "thoom" will be priceived on observing the input from alcore against a dark background. Another plan is to make a third streak of the solution on a put of prices of polished jet or black mabble, or on a plate of glass sundered at the back, and to place the streaked suitaon in front of, and at right angles to, a well-lighted window.
- The fluorescence of quantum solutions is not perceptible by gas-light, but may be brought out by burning a pince of magnesium inthon in the proper position. The use of blue glass, which transmits the ultra valid rays which produce the fluorescence of quantum, while excluding the less terrangulus any, is sometimes recommended. In this case, the light transmitted by the glass should be concentrated by means of a lens.

alcohol, but maoluble in ether or chloroform. In more dilute legudas, even if the proportion of quimne does not exceed 1 in 20,000, a deep green coloration is produced. If the green ammonicaed solution be just insuthised with acid, a blue coloration is obtained, and on adding more send a colour ranging from violet to real, but changing to green again on adding excess of ammonia

If Trim ble has proposed to use this reaction for the approximate colormetic determination of quinno. He dissolves of gramme of a quinne sait in 5 cc of fresh chlorus-water, and adds 10 cc of ammons solution. The sample is treated in the same way, and the proportion of quinne ascertained from the relative volumes of the liquid when colorined equally intensely,

The thallenoquin reaction is also given by quinidine, cupieine, hydroquinine, hydroquinine and diquinicine, but not by quinimine, or einchonine and its isomers. It is prevented by morphine.

- 4 If, after the addition of chlorine or biomine water, the quantie solution be tested with a few drops of solution of potasima ferro or ferri-cyanide, ammons being subsequently added, a red coloration is produced instead of a green. The reaction is not ochiente as the thallocquim test, but affolds useful confirmatory evidence of the presence of quinne. A Vogel modifies the test by adding bronner-water and potasissum ferrogamide to the solution to be tested, and then shaking with a fragment of marble, which, in presence of quinne, is at once covered with a red film Strychime, euchdomie, and caffeine do not give similar reactions.
- 5 On adding a fixed alkali, alkaline carbonate, or ammonia to a solution of a salt of quinine, a bulky white precipitate of the free alkaloid (more or less hydrated) is produced. The precipitate is very sparingly soluble in cold water or excess of these precipitants, with the exception of ammonia. The precipitate cannot be conveniently filtered off, washed, and weighed, as it is not wholly insoluble, and melts with very slight increase of temperature Its state of hydration is also very uncertain. But, if the liquid containing the precipitated alkaloid he agitated with ether or chloroform, or a mixture of the two, the quinine passes readily and completely into solution, and may be obtained in the solid state by evaporating the solvent. The process is readily made quantitative by operating with care and repeating the agitation with the solvent, and the quinine may be weighed in the anhydrous state as ConHaNoOo, after being dried at 100° C till constant in weight, or after exposure for fifteen or twenty minutes to a temperature of 120° C. The determination of quinine in this manner is capable of yielding very accurate results, and is of very extensive and rapid application

- 6. When quinine exists in a free state, as it is obtained in process 5 by the evaporation of its solution in ether or chloroform, it may be determined by titration with standard acid Each 1 cc of decinormal sulphuric acid (=49 grammes of H.SO, per litre) corresponds to 0324 gramme of anhydrous gumine The process is conducted by dissolving the other-residue in hot alcohol, adding as much water as can be used without causing precipitation, and titrating with decinormal acid. The indicator may be litmus, but methyl-orange or cochineal is decidedly preferable. Sharp readings are obtainable, but extreme care is necessary, owing to the very high combining-weight of quinine (ConHo, NoOo = 324). When methyl-orange is employed, the alkaloid may be conveniently used in ethereal solution, and in this case previous evaporation, as described under 5, is unnecessary, provided the othereal solution be washed with water till the aqueous liquid gives no pink coloration with phenolohthalein 1 The titiation by standard acid, of course, merely indicates the total alkaloid present, in terms of quinme The process furnishes a very useful check on the determination from the weight of the chloroform or ether-residue, and brings the alkaloid into a convenient form for further examination by one of the following processes -
- 7 On adding uncture of solme to a solution of acid sulphate of quinne in dutie alcohol, a curous compound is produced, called, after its discoverer. Here pathite, and having the formula degadaga, os 18,80,28HI, + Sang * Thus body, called also the lode-sulphate of quinine or sulphate of idequinine, is the type of a series of simular bodies formed by the action of soldine on the sulphates of the cumchona bases Herepathite is but little soluble in sold water or diute alcohol, and requires 1000 parts of hot water for solution, but it dissolves in boiling restified spirit, and is deposited on cooling in tabular crystals, remarkable for their dichrossm and their schon light,
- As quimne has no action on phenolphthalen, by the combaned use of this midiator and methy-onanges it may be determined in its salts. Standard T₀ harptis-rate ris added to the aqueous liquid until the change of the liquid copies or horse shows that the free and in mentilande. More buryts as the added slowly, with constant starring, till the prediction of a puls colour shows that the whole of the said in constantiation with the utilized in mentilsical Each 1 c. of additional T₀ stellar inequants improved the constantiation of the constanti
- ² Herepathite may be readily prepared by dissolving the sulphate of quinme in 10 parts of proof spirit containing 5 per cent of sulphune such, and adding an alcoholic solution of fodine as long as a black precipitate is produced. The precipitate is filtered off, weaked, and recrystallised from hot alcohol.

a thun film of herepathite polarising the transmitted light as completely as the tournaline. Herepathite is re-converted into sulphate of quinine by treatment with sulphurous acid, thiosulphates, sulphuretted hydrogen, and other reducing agents

Iodosulphate of quinne possesses far less solubility than the corresponding compounds of the other cinchona bases. This fact has been utilised by J E de Vrij for the determination of quinne (Pharm Joss., [3], vi 461)

With the pure alkaloid the method is capable of yielding toleably accurate results if a correction for solubility be applied, but investigations by A Christensen, B Y Shimoyama and others have shown the process to have a limited practical value as it is senously invalidated by the presence of unchondunta (Planm Jour, 13), xii 441, 1016, xxii 205, xxii 654) De Vrij's most recent method of operating is described in page 456

E B Stusrt (Pham Jour, [3], xm 1016) finds the hearpathite reaction equally delates with the thallesquit aset, and quite as easy of appheaton. The salt of quinne should be dissolved in dulte alcohol, and dulte sulphures soci, the presence of which is essential, added. Very dulte tancture of todine is then added, drop by drop, with constant agistation, when the precipitate saiddenly

¹ B. Y Shimoyama (*Pharm Jour*, [3], xvi 205) gives the following figures for the solubility of quinine heispathite in 99 per cent. alcohol at different temperatures —

	Temperature , 'O	Alcohol without Acid	Acidulated Alcohol
	15	1 in 800 parts	1 in 255 parts
	16	,, 841 ,,	
	17		1 in 117 parts.
	18		,, 101 ,,
	20	1 m 783 parts.	
ı	25	,, 660 ,,	
ı	90	,, 638 ,,	
ч			

The solubilities of the indesulphates of the principal einchona alkaloids in saidulated alcohol at 15° C were found to be as follow —

Alkaloid	Solubristy	Percentage of Iodina		
Quinine herepatilite,	1 in 255 parts	32 87		
Cinchouldine,	,, 92 ,,	63 68		
Quinidine,	,, 61 ,,	42 70		
Cinchonne,	,, 42 ,,	21 90		

appears and quickly subsides. Precipitation as herepathite may be used with advantage for separating quinine from morphine even when the relative proportions are as 1.1000

 In 1862. André (Jour. de Phasm, xh 341) described a method of estimating quinine and separating it from other cinchona bases by precipitation as the chromate, which is stated to be soluble in 160 parts of boiling water or 2400 of water at 15° C, and not hable to alteration by light or on boiling an aqueous solution A method of assaying quinine, based on the same principle, was described in 1887 by J E de Vrii (Aich, Pharm., [3], xxiv. 1073), who attributes to the precipitate the formula (C20H24N2O2)2H2CrO4, and states that it is soluble in 2733 parts of water at 12°, or 2000 parts at 16° C He directs that 5 grammes of quinine sulphate should be dissolved in 500 c.c. of hot water, and a solution of 1 2 gramme of neutral potassium chromate in a little warm water added. After standing in the cold for twelve hours, the precipitate is filtered off, washed with cold water, and weighed after drying in the air A correction of 0 005 gramme is made for every 10 cc of mother-honor and wash water. This method has been severely criticised by O Hease (Pharm Jour, [3], xvii 585, 665, xviii 582), who finds the precipitated chromate of quinine to contain 2 aqua, which fact accounts for some experimenters, working according to de Vrij's directions, having obtained an apparent excess of quinne. On the other hand, canchonidine and hydroquinine are in part thrown down with the quinine, which renders the method inapplicable for separating quinine from its most constant associates

parating quintne from its most constant associat

Quintne is distinguished:---

1 From cinchonine, a, by its fluorescence, h, its lavorotation, c, the thallenoquin test, a, the crystallisation of the sulphate, e, its solubhity in ether; f, its solubhity in animonia, g, the sparing solubhity of the noiseulphato

2 From cinchonidine by most of the above leactions, except b, and less sharply than einchonine by those tests depending on relative solubility (\hat{d}, e, f, g)

3 From quinidine by b, d, f, q, also by (h) yielding no precipitate with potassium iodide, and (i) the insolubility of the sulphate in chloroform

4 From quinamine by b, c, j, proceptation as tartrate; and k, the sparing solubility of the sulphate

5 From cupreine by a, and (i) the insolubility of the precipitated alkaloid in excess of soda

Methods for the separation of quinine from the associated cinchona bases are given on pages 411, 453, et seg The separation of quinne from morphisms may be effected, as already stated (page 405), by precipitation as herepathite, also by treating the free alkalouds with chloroform or ether, which leaves the morphism undissolved

From strychnine, quinine may be separated as indicated under "Easton's syrup" (page 377)

SALTS OF QUINTNE,

Quinne is a strong base, completely neutralising acids, and forming crystallisable saits having no reaction on lituus or methyloronge. These saits react with phenolphthalam as if the acid were in an uncombined state. Quinne also forms a series of acid saits, of which the acid subplate of quinnes is the type.

Several of the salts of quinine are official in the Pharmacoposia,

and others are extensively used in medicine

Quimine Sulphate. Diquinic sulphate $(C_{20}H_{24}N_{2}O_{2})_{2}H_{2}SO_{1}$. This important salt, sometaues called "disulphate" or "basic as uphate" of quinine, forms, in the hydiated state, the ordinary medicinal sulphate of quinine of commerce

Sulphate of quinine is usually met with in exceedingly light scales, or long, flexible filtium neodles, having a macreous aspect and a pure and intensely bitter taste

The crystallised sulphate of quunne of commerce usually contants about 145 per cent of water, a proportion which corresponds closely to a 7-atom hydrate, which requires 145 per cent, According to some authorities, however, the wholly unefforesced crystals contain 8 aque, or 4 any 145 7 8 aqua * H B, Parsons

¹ Chemically rus optimus sulphate, free from hydrocytaline, crystalizes in heavy needles resulting sulphate of mer. The high chances of the commercial sails a cheefy due to the presence of small edunitaries of the anjohase of hydrocytanon and unchoundary, and possibly of hydrocytalonolary of high product of the heate of the sunchonial of hydrocytalonolary of

² The British Pharmacopeas of 1885 gives the founded of crystallised quante sulphate as (B.H.SO.), 15H.2O, which corresponds to 7½ aque. The freshly prepared self is stated to lose 15 2 per cent. of water when dried at the temperature of boiling water.

(Proc. Amer Phasm. xxxii 457) has published the results of drying for three hours, at 100°, 1015 samples of gumine sulphate (taken from time holding 100 ounces each, and not previously opened) of American, German, and Italian manufacture. The average loss of water was 13 84 per cent, the highest average from any one maker being 14 36 per cent A. J. Cownley (Pharm. Jour, [3], xvi 797) found the water in thuty-seven samples of commercial quinine sulphate examined during the two years prior to 1886 to range from 810 to 1612 per cent D Hooper states that the water ranges from 5 to 18 per cent Hesse (Ber. xm. 1517) states that pure crystallised guinine sulphate, which has not effloresced, contains 8H2O, or 1617 per cent of water. Cinchoniding sulphate, on the contrary, crystallises with 6H.O. or 13 7 per cent Hence, if a sample of gumine sulphate be dry and quite free from efflorescence, the proportion of water is an indication of its number

Crystallised quame sulphate is readered perfectly analysious by exposure to a temperature of 100° C. If a laghet comportance be supployed for its delaydration, there is a danger of some of the alkaled undergoing convenient into quamens (see penge 43). If the anhydrous sulphate of quame he exposed to most arr, it is particularly absorbs from 48 to 5 per cent of water, a proportion which corresponds to the formula B.H.S.Q.+2HIQ.O¹ On the other hand, the crystallized and it rapidly loses water on exposure to other hand, the crystallized and translation of the factor which are the arm of the control of the decimal points of the decimal points are decimally decimally decimally decimal to the decimal points. The same quantity of water is retained when the crystallized can strong alcohol

Quinnes sulphate requires 750 parts of cold water for solution, but desolves in about 30 parts of water at 100° 0. It is far less solution in water containing sulphate of magnesume, sodium, or aumonum than in pure water. In a strong solution of Rochelle silt, quinine sulphate as of little soluble that the alkaloid can scarcely he detected by the fluorescence or thallstoquin test. On the office land, the solublity of sulphate of quinnes water is increased by the presence of ammounum chloride, or of potassium intuites of chlorides

In alcohol, quamne sulphato dissolves more readily than in water, requiring only 7 or 8 parts at a boiling temperature, but is much less soluble in cold spurit (see "Inneture of Quamne," page 423) Quanine sulphate dissolves in about 24 parts of cold glycern, the solution bung precipitated by addition of cold glycern, the solution bung precipitated by addition of water Crystallised quamne sulphate is not soluble in fixed oils.

¹ H P Parsons recommends the official adoption of this hydrate as a definite and stable form of quinine sulphate

ether, chloroform, or petroleum spirit (It is said to dissolve in benzenc.) In the anhydrous state, I part of quinine sulphate is soluble in about 1000 parts of chloroform (see page 416)

In dulue sulpiuros and, quanno sulphate is readily soluble, owing to the foundson of acad sulphate of quanue, $C_{\rm BH}^{-1}$ N₀, $C_{\rm BH}^{-1}$ The crystallised sail bases 6 again in the existencing TH₂O. The analyticus at 100° C. When heated to about 136° C it melts, and is convexited into the corresponding compound of quanticus (see a gage 434). And sulphate of quantum devices in 11 parts of cold what, and more readily in hot water or in alcohol to strongly fluoressents adultions

From a solution of quinne in excess of dilute sulphunic and, an acid sulphate may be obtained, having the composition $C_{10}H_{24}N_2O_{2}2H_2SO_{4} + 7H_2O$ ($= C_{20}H_{24}N_2O_{2}H_2SO_{4} + H_2SO_{4} + 7H_2O$

Normal quinne sulphate has a specific rotation in alcoholic solution of S_s = 1915°, alculated for the anhydrous salt. Excess of and increases the rotatory power. When dissolved in water activatived with hydrochloric acid, the value of S_s at 15° is 233°75° (If c s s c).

Sulphate of quinme is largely employed as a febrifuge and tonic, the official doso ranging from 1 to 10 grains. It has marked antiseptic properties.

The fluorescence of sulphate of quinme is considered on page 400, its reaction with indune on page 403, and with the thalleroun test on page 401

Examination of Commercial Quinine Sulphate

The salts of quame, except the tanaate (page 420), can all be examined by the following methods applicable to the sulphate of quame, provided they are first treated with 10 parts of boiling water and their own weight of sodium sulphate. The sulphate of quame which deposits on cooling and the mother-liquor obtained can then be examined in the usual way

Commercial sulphate of quinnia was formerly subject to adultorions of a very gross chancter. Among the bodies amployed to sophistoste it are said to have been starch, gum, storain, sahen, phloridam, sugan, sulphate of magnesum, sulphate of sodium, achalic, sebestos, bout each, &c. Some of these additions are sportyphal and the majority are certainly obsolete

Mineral additions would be readily recognised on igniting the sample, which, when pure, will leave no sensible ash. Starch, chalk, stearm, and boric acid would remain insoluble on treating the substance with cold dilute sulphuric acid, and gum would be proceptated on adding excess of alcohol to the solution thus obtained. Soluble impurities generally may be detected and estimated by dissolving the sample in het water and adding excess of haryta water. The alkaloud is then removed by agitation whether. After removing the ethereal layer, a stream of carbonic and its passed through the aqueous liquid to prequisite the excess of baryts, and the whole wall boiled and fillered. Sulphinte and carbonate of barum will be left insoluble, and the fillente will contain any sugar or other soluble impurity present in the original sample, and the observation of the weight of the renduce left on evaporation will allow of a determination of the amount. In presence of sugar the liquid will excit a dextro-rotatory action, and in presence of educar is layer-ortatory action, and in presence of educar is alway-rotatory action, and

Treatment of the original solid sample with concentrated sulphure and, attended by gentle warming, will suffice for the qualitative detection of some imputities. Sugar and maintet will become charred, while sahun developes a striking red colour Good commercial quinne sulphate dissolves with fami yellow colour in strong sulphure and, and the tent is not deepened on

warming gently

Similar general impurities may be rapidly tested for by a tost devised by He sae, and discerbed on page 417 Solicus, if present in greater proportion than 1 per cent, may be detected by this test. The reaction mothation with collection of the test of the reaction mothation will be coloured deep red by concentrated sulphurne and, and will reduce Felling's solution after boiling with children sulphurne and the reaction with strong sulphurne and will be produced by the original sample if the proportion of salicus has provided by the original sample if the proportion of salicus may be detected in the filtrate from the prespitate produced by adding baryta to the exposure solution of the sample Another test for salicus is to dissolve 0.25 gramme of the sample if salicus be present, on boiling the highly for some minutes a white turbulity will be produced, due to the formation of salire-time.

Sulphate of quaine has occasionally been largely adulterated with or entirely substituted by the hydrochlorule of cunchonne. This fraud is recognisable by testing for chlorides with intin acid and mutate of silver, and for cinchonne as described on page 413

The most common unpurity of commercial sulphate of quante as an admixture of one or mose of the sulphates of other concloraallatoids, especially cinchonidine. This admixture is often purely seedental, owing to imperfect separation of the other alkaloids during manification, but is no doubt semetimes provided for and secured by suitable arrangements of the manufacturing operations, while occasionally an intentional admixture of other alkaloids has occurred

Manutachurers of quinnes sulphate produce at least four qualities of the article (1) The pure salt or "heavy sulphate," of which the use has been hitherto extremely limited, chiefly on account of its unfaminarity to the membors of the medical profession, (2 and 3) products estadying the requirements of the German and Datch Phermacopenase, and (4) products satisfactly others than the above-mentioned pharmacopenas and containing from 4 to 6 per cent of sulphate of unchonduler. Other products may have a certain commercial importance, but have no "legal status" in cuvised countines.

The best samples of commercial quinne sulphate are soldom free from enchomdine, but contain not more than 2 or 3 per cent, whilst other kinds contain from 5 to 10, and even 20 per cent, of enchonidne sulphate, and on one occasion B H Paul found 60 per cent.

F W Fletcher (1882) states that quinme of English manufacture is usually practically free from cinchonidine, but that certain foreign brands always contain from 10 to 15 per cent, in one case the proportion exceeding 25 per cent A. J. Cownley has published determinations of circhondline made by a reliable process, and finds the proportion to range from nil to 139 per cent , the next largest amount being 9 0 per cent. More iccently (1889), Paul and Cownley (Pharm Jon., [3], xix 665) found the cinchonidine sulphate present in twenty-three typical samples of quinine sulphate, representing all the different makers. to range from nil (in two instances) to 12 34 per cent. In fourteen out of the twenty-three the proportion of impurity was less than 6 per cent The two samples which were wholly free from cinchonidine were probably manufactured from curies bark, which is characterised by the absence of cinchonidine, and in one instance this conjecture was confirmed by the detection of a trace of cupreine in the sample. In addition, hydrogumne is a very constant impurity in quinine sulphate, a very notable proportion being sometimes present, and, according to Hesse, hydrocrachonidine and homocinchonidine may also be met with in quinine from certain sources The presence of even 1 per cent of cinchonine or quinadine in quinine sulphate is far more likely to be intentional than due merely to accident or careless manufacture, but these alkaloids are apt to be met as accidental impurities in quinine hydrochloride.

The detection and estimation of foreign alkaloids in commercial

sulphate of quinine has received much attention, and considerable ingenuity has been exercised in the solution of this somewhat difficult problem

The recognized methods of testing commercial quime sulphate for admixtures of other alkaloids are, for the most part, based on the removal of the greater part of the quime as a spungly soluble sulphate, and the datention of the remaining quime from its associates by its greater solubility in other and its sububility in excess of aminoma. A great variety of tests based on these principles have been devised, especially for recognition and estimation of cinculditie, the detection and determination of the other alkaloids when present in notable proportion presenting companiatively little difficulty

The separation of small proportions of cinchonduc from comme is particularly troublesome, and formerly any considerable propertions of the admixture must have escaped recognition B H Paul (Pharm Joss, [3], vii 653) has shown that when the test for quinine sulphate prescribed in the Bistish Pharmacopasis of 1867 is rigidly adhered to, it is difficult to detect an admixture of 20 per cent of the curchondine salt. By reducing the volume of ether used, any impurity in excess of 10 per cent may be detected, but less than this proportion escapes recognition, owing to the property possessed by quinine of increasing the solubility of cinchonidine in other, or at any late of preventing the latter from separating in a crystalline state. Hence, for the detection of small proportions of canchonidine, it is necessary first to separate the greater part of the quimine. This may be done by utilising the fact that quinine sulphate requires about 750 parts of cold water for solution, while curchomdine sulphate is soluble in 100 parts. This principle was originally employed by Kerner, but its application has been modified and improved in several respects by Paul (Pharm Jour, [3], vii 653, xvii 645), and Hesse (Pharm. Jour , [3], xvii 975). But cold water does not completely dissolve emchanidine sulphate from commercial quinine sulphate, according to Hesse, because of its existence in the form of a double sulphate of the two alkaloids This compound is decomposed or disintegrated by hot water, even if the quantity of liquid be insufficient for its solution, the cinchonidine salt passing almost wholly into solution, while the quinme sulphate is for the most part undissolved. On the point whether it is better to treat the sample with water at 60° or to 100° C, authorities are at variance Hesse considers that at a boiling heat more of the quinine sulphate will pass into solution, and hence there will be a greater tendency to the re-formation of the double salt when crystallisation takes

place. Kerner and Weller also recommend the use of water at 60°. E Jung flexes h (Jour Pharm et Olium, [5], xv. 5, Pharm, Jour, [5], xv. 1585 gives the preference to a boiling temperature, and points out the tendency to erratic results it less heat be employed Paul (Pharm. Jour, [3], xvn 595) considers that the best results can only be obtained by using meally sufficient water to effect the complete solution of the quinne subhate at the boiline-round

The mode of operating recommended by Hesse is to take I gramme of cumine sulphate previously dried at 100°, shake it with 20 cc of water at 60° C, filter after cooling, and agricate 5 cc of the filtrate in a narrow tube with 1 cc of other and 5 drops of ammoma (sp gr 0 96) The clear ethercal solution thus obtained should not deposit crystals on standing. If, on leaving the tube at rest and in a closed condition for two hours. the ethereal stratum be found free from crystals, the sample may be considered pure , but if it contain more than 0 25 per cent, of cinchonine sulphate, 0 5 of quinidine sulphate, or 1 0 per cent of canchonidine or homocinchonidine sulphate, a distinct separation of crystals will occur. The last two impurities appear granular, while crystals of cinchonine and quinidine form concentric groups of delicate needles If the proportion of cinchonidine be as high as 3 per cent, the separation of crystals will occur immediately, or within three minutes, 2 per cent will show in about ten minutes, while with less than 1 per cent no separation will occur even after twelve hours 1 To detect smaller proportions of these alkaloids, the cork of the tube should be replaced by a loose plur of cotton-wool, so that the ether may gradually evaporate On examining the residue with a lens it will appear distinctly crystalline if 1 per cent of cinchonidine or homocinchonidine sulphate be present, and a mere trace will be recognisable by the presence of a few crystals m the amorphous mass of quinine 05 per cent of cinchonine sulphate, or 10 per cent of quinidine sulphate, will cause an almost immediate separation of crystals from the other. Their presence is far more likely to be intentional than merely accidental or due to careless manufacture.

The British Pharmaroposia of 1885 gives the following methods of testing commercial sulphate of quinine for accompanying

A deport of cinchondine is recognised by the capillary mang of the precription beyond the othercal layer numediately after shaking the solution. With a large proportion of cinchondine a white chalky mig appears at the line of contact of the two lumbs.

- alkaloids 1 The salt "should not contain much more than 5 per
- a Test for Conchantding and Curchange Heat 100 growns of the sample in 5 or 6 owners of boiling water, with 3 or 4 dions of dilute sulphune and 2 Set the solution asule until cold Separate by filtration the numbed crystals of gumme sulphate which crystellise out To the filtrate, which should nearly fill a bottle or flask, add other, shakme occasionally, until a distinct layer of ather remains undissolved. Then add ammonia in very slight excess, and shake thoroughly, so that the quimne at first procontated shall be redissolved by the other. Close the flask, and allow it to stand for some hours, and then remove, with a pipette. the supernatant, clear, ethereal layer which should occupy the neck of the flask. Agitute the residual aqueous hould and the separated crystals of alkaloid once or twice with a very little ether Collect the separated alkaloid on a tared filter, wash it with a little ether, dry at 100° C., and weigh Four parts of the product represent five of crystallised subpliete of cinchonna or curchondine in the samule
- b Test for Cuprence Shakes the crystallised sulphate of quames obtained in Test or with I fluid onne of either and \(\frac{1}{2}\) fluid quames of ammonia (sp. gr. 0.959), separate the ethereal solution, and add to it the ethereal solution and westings obtained in Test a Shakes the united othereal liquid with 1\(\frac{1}{2}\) fluid onne of causties sold solution (10 per cent), adding water if any solid matter separates Separate the ethereal layer, agatate the aqueous liquid with more ether, and separate as before Heat the squoous liquid with more either, and separate as before Heat the squoous liquid to boiling, and exactly neutralise is with dialte sulphine and Allow the solution to cool, separate any crystalline cupreme sulphiate by a taref filter, wash with a title cold water, thy and wear?
 - c Test for Quantitine. Recrystallise 50 grains of the sample as
- 1 The Procal Colors of 1384, making two of Konne's method of analyses, or presention that 5 o. of a motive hope obtained at 15° O, aften teatment of 1 gramme of the officiant sailt with 10 c. of linke warm water, shall resumment profestly implied for 24 hours after the addition of 7 c. of a whitten of ammonia of 0 69 specific gravity. The manufacturian considered these neglections server. However, the new American Mehromeopora practice the time of 7 c. of ammonia, which notly slightly less servers a test, and the plannic operate of Russey, Finland, Sweden, the United States, and Japan have adopted nearly the same test. The Dubit Pharmacopeas has related to 4 ce or mount of ammonia to 5 c.a., and the German Pharmacopeas of 1350 to 4 ce.
- ² This addition of sulphuro and is objectionable, as tending to moreave the solubility of the quainte sulphate and diminish the deleacy of the text. It would be better to direct the addition of just sufficient and to lender the solution family send to himne.

just described in Test a, and to the filtrate add a strong solution of potessum noble, and a little netified sparit to prevent the precipitation of the hydroddes of amorphous bases Collect the precipitate of quantine hydrodded, wash it with a little other precipitate of quantine hydrodded, wash it with a little other wash, and the precipitation of the precipitation of the analysis of the precipitation of the precipitation of the serial vegets to design and the serial vegets to design and the serial vegets of the precipitation of the serial vegets of serial vegets serial serial vegets serial serial vegets serial serial vegets serial serial vegets se

The foregoing tests ane, of course, not intended for the detection attends of astimation of minute traces of accompanying alkaloids in quinne sulphate. Conclouding his about one-fourth the potency of quinne, and hence the therapeutic value of the preparation is not so greatly affected by a small admixture as is the commercial value.

B H Paul (Pharm Jour, [3], xvn 647) points out that the deheacy of the test would be much increased by evaporating the filtered aqueous solution to about one-fifth of its volume before shaking with ether and ammonia 1 Operating in this manner, as small a proportion as I per cent of emchandine sulphate can be detected with certainty, even when only 10 grains of the sample is employed, provided that the closed tube (employed with small quantities instead of a flask) be allowed to stand for at least twelve hours for the formation of the crystals De Vrij (Chem Cent) , 1885, 968) has suggested the addition of sufficient sulphuric acid to convert the bases into acid salts before separating them by finetional solution and crystallisation. Hesse (Pharm Jour, [8], xvii 486), who expresses a high opinion of this method if calefully performed, recommends the following mode of operating -5 grammes weight of the sample is dissolved by the aid of heat in 12 cc of normal sulphure acid (49 grammes HoSO, per little) contained in a small porcelam basin, and the solution poured into a funnel closed at the bottom,2 in which it is allowed to cool. At the end of two hours crystallisation is complete, the stopper is removed, and the mother-liquor allowed to drain away as completely as possible, its removal being assisted by suction. The upper portion of the crystals is then pressed down with a glass rod and washed with 3 cc of cold water, added drop by drop while the suction is kept up The whole solution is then mixed with 16 cc of ether (so gr 0 721 to 0 728) and shaken up 5 Three cc of ammonia (sp.

¹ In a later paper (Phorm Joss , [3], xx 665) Faul and Cownley recommond that the solution should be concentrated to about I fluid drachm (3½ c.o.), and the deposited crystals separated before treatment with summons and chiral fluid flui

This may be conveniently effected by a glass rod introduced from above, and having the lower and covered with a short length of india-rubber tubing. The same rod can be afterwards used for pressing down the crystals.

S If the sample contain more than 10 per cent, of emchanding the volume of ether must be increased.

gg 0.960) is next added, and the whole well shaken again. After stanning one day the other is removed with a pipetic, and the crystals which have separated are collosed on a liker and weshed with water stantand with teher. The liker is then placed on a absorbont surface, the crystals again washed with some either and dand at 100°. These crystals are not pure carehoustune, but a compound of quinne and emchandine, having the composition $C_{\rm HB}N_{\rm Q} > 26_{\rm HB}N_{\rm Q} > There is always a certain natural,$ in the ample is very small, and hence Hesse recommends that theweight obtained should be multiplied by 0.62, instead of by 0.615,which is the cicculated factor for the above to mind.

B. H Paul (Pharm Jour., [3], XVII 555) strongly objects to the acid sulphate test, on the ground that the crystals of acid sulphate are not free from cinchonidine, while the amount of quaine retained in solution is so much increased as to interfere with the subsequent crystallisation of the enuchonidine from ether

Conversion of quinne into and crystallisation as the acid sulphate effects a separation of hydroquinne, which remains in the motherleques, while repeated recrystallisation of the neutral sulphate tails to effect this (compare page 424).

A mothod of assaying quinine sulphate for einchonidine, based upon the optical rotation of the solution, has been recommended by several emment authorities and is equally distrusted by others Oudemans was among the first to experiment in this direction, and Hesse proposed a definito process of assay, based on the rotation of the sulphate Koppeschaar proposed to employ the tartratos by preference, while R H Davies operated on the sulphates De Vrij has strongly recommended the optical method of examination, giving preference to the taitrates Jungfleisch and Paul and Cownley have expressed strong distrust of the optical method, considering it manifestly impracticable to determine proportions of 1 and 11 per cent of cinchonidine in quininc sulphate containing even minute proportions of the cinchonine and quinidine salts, and D II o ward states that no published method gives the mixed testsates of quinine and cinchonidine sufficiently pure to render the polarimetric assay absolutely rehable. Hesse has modified his former high opinion of the method, and points out that it is invalidated by the presence of hydroquinine, which is invariably present in commercial quimine sulphate, and is not separated by converting the bases into tartrates.

¹ Hesse's test-experiments on mixtures of pure quinine and einchonding sulphates in known proportions justify this empirical factor.

The presence of I per cent of hydroquames sulphate roduces the rotation to the same extent as 0.42 per cent of the unchondine salt, and its presence accounts for the excessive and theoretical figures for candomium often obtained by those who rely on the optical method of assay. Hydroquame cannot be perfectly separated from quame even by repeated recrystallisations of the north sulphate, but it can be completely got rid of by converting the alished in the is each sulphate and isopratising this from water or alcohol, when the hydroquame remains in the mother-liquer (commerce ance 424)

compare page 43*)
For the optical assay, Koppeschaar (Zeitsch Anal Ohen, xxv 362) recommends that the quime and cinchondrine should be convested into tatrates by pacequisting the neutral solution with Rochelle sait, and the precipitate washed with a hittle cold water and duot at 135*-130* °C; 0 400 gramme of the dry product is then dissolved in 3 cc of normal hydrochlore and, and the solution diluted with water at 15° C to a volume of 20 cc.
The solution is placed in a packeted tube kept at 15° C, and the totalory power observed by a polarimeter employing monochromatic solution. The solution is placed in a packeted tube kept at 15° C, and the totalory power observed by a polarimeter employing monochromatic solution.

where S is the specific and a the angular rotation, and l the length of the tube in decimetres. From the figure thus obtained, the percentage of quinne tartiate, z, in the mixed tailable may be ascertained by the following (Koppeschaar's) formula

$$x = \frac{100(8-13767)}{824}$$

Each 1° of diminution in the specific rotation below 220 07° coreposed so bout 1 2 per cent of cuchonvilue attratus in the mixed tartines. The angular rotation is diminushed by 0.077° only by the presence of 1 per cent of cuchondules tartinet. Notivithe standing the extreme accuracy of observation necessary, H oo per $(Pknam\ Jow)$, (3) xvii (1) has found the opheal detarmination of quinno in the mixed tailtastes to give very satisfactory rosults. It is as found that epocific totation of quinne, hydroquinne, and cunclosulture tartiates, for Oudemans' concentration Π_i to be respectively, $-2.12.5^\circ$, -176° , and -13.2° , -176° , and -13.2° .

For the detection of conchonnel or quantities in quantite sulphate, Hesse proposes to dry the salt at 100° C, and agitate 1 gramme with 15 cc of chloroform free from alcohol Tho liquid is passed

According to Laborde (Phasm Joss, [3], xm 684) the presence of ounchounce materially alters the physiological effects of quinine salts.

through a small filter. If 10 c., on evaporation at a gentle heat, leave an anorphous readies weighing more than 035 gamma, euchomore or quindine sulphato is certainly present. If the resulue be crystalline and less than the above weight, it may be tested for the foreign alkaloids by heating it with 5 c. of water, adding 2 gramme of potassaum sodium tartasis, cooling, filtering from the precipitated quinne and emohandine tartasis, and mixing the littate with an equal volume of ammonia. If quindine or cuchosine be present, a precipitate will be formed, and may be further examined by agriation with other (see page 412), or by treatment with tochie of potassum (see page 413). Sulphate of emchanding if present, will assum undissolved by the chloroform, but will swell up into very bulky needles, which suck up the chloroform, but will swell up into very bulky needles, which suck up the section of the first as somes and do not viged it assum without pressure

L Schafer (Arch. Pharm., [3], xxv 64, 1033) has described a method of testing commercial quimine sulphate, based upon the precipitation of the boiling aqueous solution by neutral potassium oxilate. After cooling and filtering, the filtate is tested by

addition of caustic soda.

O Sehliekum (Arch Phann, [3], xxv 128) has myestigatal De Yriy's chromate nethod (page 405), and funds it applicable, under certain conditions, to the examination of quinne sulphate. On preceptating a solution of this or other notical quinne salt with neutral potassium chromate, and filtering after four or more hours, the filtrate remains clear on addition of soda, if the quinne salt was pure. In presence of \(\frac{1}{2}\) per eent of cunchonine sulphate, or 1 per cent of the quinndine or enchonidine salt, a turbulity is produced at once or after a time

A test for the purity of quinne sulphate, devised by H esse and adopted by the German Pharmacoppenia, consist in Instantage I gramme of the sample for a short time to 40°-50° C, in 7 e o. of a mixture of 2 volumes of abloroform and 1 of absolute about 10 H is sample be pure it is completely dissolved, and the solution remains quite clear on cooling Sulphates of other conchons bases and varous organic and morganic impurities remain unsabble (compare page 409).

A somewhat similar test has been described by E H 1: s e h s o h n, according to which 0.2 gramme of the quinne sulphate should brinkly agatated with 5 e o of a nixture of 30 paits of patroleum elsher of 0.680 pg gr, with 70 parts of abloroform. The liquid is hiltered, and diluted with three or four times its volume of petroleum elsher, when an admixture of 0.1 per cent, of sulphates of oblice enchons bases will give rise to a turbedity or picunisate

For the detection of amorphous alkaloid in commercial quinine

sulphate, De Vrij recommenda the following method: —The sample is discolved in dilute and, and shaken with ammonia and either for estimation of total alkaloud. Sufficient decincemal oxale and is alided to the ether-resolute to convert the alkaloud into nontrall oxalets, and the higud is evaporated at a steam-heat and the residue thoroughly drad in the water-lath. It is then dissolved in chloroform, and the higud filtered if necessary. The clear solution is next teated in a test-tibe with a few drops of west, when crystals of oxalete of quinne will appear in the chloroform. If the sample were pure the aqueous layer will remain clear and uncoloured, but if amorphous alkalot be present it will be dissolved by the water and colour it vallow.

Quantum Hydrochlorate Hydrochlorate of quimme B,HCL This salt forms long absotso-the pranse nonthining 2 aqua, which become anhydrous at 120°C without previously melting The debuylunted salt fuses at 168°-160° without change, and is not converted into quincing, as stated by Pasteur (Hesse). If an aspisous solution of quimno hydrochlorate saturated at 15°C be allowed to stand for some time at about 0°C, large outsideful as solvible in about 4°C points of quimno hydrochlorate aguinno hydrochlorate average and the proposed solving the proposed proposed to stand for some time at about 0°C, large outsideful as solvible in about 4°C points of cold water, and very solution in hot water and in about 1°C points of cold water, and very solution in hot

Quinne hydrochlorde has been frequently substituted of late years for the sparingly soluble sulphate Thus it is used in making the Tineture of Quinne, B P The hydrochlorde is the more expensive salt, Jowing to the increased difficulty of crystalhang and the high penentage of quinne contained in it (84-2 ve cent. accusate 75 5 in the crystallistic sulphiate).

Quinine hydrochlorude is propared by reacting on the sulphate with chlorude of barrium. Hence it is apt to contain either undecomposed sulphate of quinine, or else barrium chloride. The latter impurity is, of course, very objectionable.

Quunine bydrochloride may be assayed in much the same manner as the subplact (see page 408 et say). The BP test for quunner sulphate is applicable to the axammation of the hydrochloride, if the sample be purvously dissolved in ten times its weight of boding distilled water, together with its own weight of crystallised solution sulphate the crystals of quinne sulphate which are deposited, and the filtrate from them, can thon be examined as

¹ The and hydrochlor ad, EH₂(II₂, as obtained by pracipitating the acid sulphate of quinne by barnum chloride. It forms groups of concentron needles, which can be dired without change at 110°, and are soluble in an equal weight of water. It also separates as a gelatinous mass, which becomes crystaline on gentle warming.

described on page 412 et seq. The hydrochloride of quimine is more likely to be contaminated with the similar salts of einchonine and quimine than with the hydrochlorides of einchondine and homosuphonidae.

Quantine hydrochlorade has on several occasions been acadentally mixed with or replaced by the corresponding salt of morphism. The impurity may be detected by warming the salt with dilute nitric and, which acquires a yellow or red coloni if morphine be present, or the salt may be placed in a pencelain cuclide and mostened with very neutral ferric chloride, which will produce a given or blue colour if morphise be present. The production of a blue colour with mixed solutions of ferric chloride and potassium feniverantic (page 317) is also well adapted for the detection and approximate estimation of morphism in presence of cunchons lauses Lastly, the expectors solution of the salt may be treated with animonia and agetated with a small quantity of ether, when any morphism (or unchosmo) will remain undiscoption.

Gunnie Hydodomula, BIRN-H.Q. is prepared by mixing equivalent quantities of quinne sulphate and potassium bremitle with their own weight of water, adding three or four parts of strong alcohol, filtering from the preceptated potassium sulphitude and erystallising the quinne hydrobromide from the filtrat real trongs silky needles, soluble in 16 parts of water to a solution and to be fluorescent (f)

Quarine Carbonate, E.H.,CO₃+H₃O_. is obtained by passing carbon dioxide into water containing freshly precipitated quinnic hydrate, and exposing the resultant solution to the au I torms tansiticent needles, efflorescing rapidly in the air, decomposing at 110°C, and soluble in water or alcohol but insoluble in either

Quintine Chromate, B.H.G.O., +2H.O. The anhydrous selfmptidly re-absorbs 2 aqua on exposure to arr. It is soluble in about 2000 parts of cold water, and has been recommended by d. s. V111 for the determination of quinine (page 405). It becomes anhydrous at 80°7, and decomposes at a higher temperature.

Gassians Cralate, B.H.C.O.4+6H.O. forms delicate needles soluble in about 900 parts of oold water. The oxinhets of the other frequently countring cinchona bases are companitively easily soluble, and L. S ch a f or has based on this fact a nethod of separating small proportions of these bases from quinne (page 41).

Gunnare Valerate forms coloumless rhombondal plates, having a pearly lustre and a faint odour of valeric and It is not dehquescent, and fuses at a low temperature Quinne valerate requires 110 parts of cold or 40 of boiling water for solution, and is easily soluble in alcohol. Valerate of quinne is liable to con-

tain much the same impurities as the sulphate (see page 408). Sulphate and hydrochloude of quinine, and valerate and acetate of zine are also liable to be present

Questine Tannate has some into use in medicine on account of its comparatively tasteless character. The commercial product varies greatly in its composition, the bitter taste decreasing with the amount of alkaloid contained in the specimen.

For the preparation of quinne taininte, Pell's recommends the preequitation of a saturated solution of 1 part of quinne hybric chloude by 3 of stanim (in 10 per cent solution previously neutralised by ammonia). After standing treatify-time hours, the wested precipitate is draid at a low temperature. So prepased, quinne tainiate as a yellowshe-white amorphous powder, soluble in about 50 parts of cold water or alcohol. Its solution gives the rescuess of stanies can be solved to the solution of the solution

In some cases, the quinne in the commercial tannate is largely explaced by other cunchons buses. The following analyses by Jobst (Arch Pharm. [3] xm 331, Joun Chem Sec, xxxiv 678) illustrate the composition of commercial "tannate of cumns".

	1	2	3	4	5	6	7
'Water lost at 120° C	7.2	97	91	98	10 2	10 7	11 4
Qumine, Qumidme, Cinchondine, Cinchonne,	81 87	22 72	4 46 11 97 7 88	13 10	6 23 Tiace 23 80 Tiace	10 00	7 40
Total Alkaloid, .	81 37	22 72	28 76	23 82	27 08	10 00	7 40

To secretain the proportion of total alkaloid in quinnie tannies, Jobst products I grainue of the sample, and mixes it with milk of lime. The invitine is direct on the water-bath, and the resulting powder exhausted with chloroform. The chloroform is filtered, evaporated, and the resulting weighted after drying at 120° C. The alkaloid thus separated can be further examined as described on page 412. There seems no reason why the mixture of the sample with milk of lime should not be agtated directly with chloroform, thus avoiding the evaporation to dryiness of the argument should. A similar process is adopted by S. N eu mann, who agitates the finely divided tannate with strong solution of caustic alkini and excess of their. The presence of solid particles

in suspension, either in the ethereal or alkaline solution, shows that the sample is impure or that it has not been completely decomposed.

 Quante Tartiate, n₂H₂C₂H₄O₆+H₂O, forms a crystalline prequitate, soluble in 910 parts of cold and more readily in hot water It becomes anhydrous at 100°, and is the best form for observing the ordical activity of quinnie (page 416)

Citrate of Quintum is not a commercial preparation, but in combination with terms citatle it constitutes the Ferre et Quintum Citats. R.P.

Citizet of Ion and Quantus occurs in commerce in the form of thin transparent dolsquescent scales, varying in colour from a doltrate greenish golden yellow to yellowish brown, escording to the proportion of aminonium critate present. The jusquation should be somewhat slowly, but freely and completely, solublo in cold wate. It is insoluble in alcohol or other The aqueous solution has a very bitter and chalylosate tests, and should be only very slightly and. On adding aminonia to the cold solution, while quinne hydrate is thrown down, and the hydra dissumes a darker colour. No ferric hydrate is preceptated unless the hyurd be heated, or a fixed alkin slushfutured for the aminonia.

Citrate of iron and quinine is hable to several suphistications

. The proportion of water in the sample may be ascertained by drying a weighed quantity in the water-oven. It averages 8 per cent, and should not exceed 10 to 12 per cent.

Adultenation with potassic-actuate or potassac-tarriate of iron would be detected by the strongly alkaline seaton of the resultue left on agmiting the substance, a genuine preparation yielding and natural value of the thirus jupon. The substitution of tattane send for the estime send of the sample is now marpotable, but may be detected as described in Volume I.

The proportion of coals of son can be estimated in the pune preparation with sufficient accuracy by springs a known weight of the sample. After testing the sait for fixed alkalı, a few drops of mitric such should be added and the resulte again spitted. This treatment ensures the complete combustion of the carbon. Citrate of iron and quame ought to yield from 18 to 20 per carbon of fenic oxide on spittin. A more accurate estimation of the iron can be made in the sait, if desired.

Excess of citive acid is indicated by the extra acidity of the sample, but the commercial substance frequently contains a much larger proportion of acid than is prescribed in the British Phanmaconomia.

Sulphates are almost invariably present in citrate of iron and

quinne, owing to imperfect washing of the ferne hydrate suployed, or to the introduction of the quinne as sulphate nation of preespitated hydrate. The employment of sulphate of quinnes is said to render the preparation liable to yield a turbul solution, but it has the advantage of preventing the moviable loss of alkaloid attending the neparation of quinne hydrates.

The British Pharmacopenia of 1867 required that the crimic of tron and quinne should contain 16 per cent, of alkaloxia, as determined by drying, at an unstated temperature, the unwashed quinner hydrate procupitated by ammoun. In the edition of 1885, this faulty process was substanted by a method recommended by the substance of the process of the substance of the quinner from the aqueous solution by ammouna and extraction of the calknoid by ether or chloroform. No temperature is prescribed for drying the alkaloxial resolute, but a constant weight is best obtained at 110°-120°. By this process, which yields very accurate region to the preparation is now required to yield 16 per cent, of alkaloid ² If preferred, the resultion may be dissolved in a little solo, this solution diluted with water, and trinsted with a standard numeral seld and methyl-coarse.

The proportion of alkaloid in the citiate of iron and quinine of commorce is often notably less than the 15 per cent required by the British Pharmacopava (see Phana, Joan, xvii 234, xix 259, xx 1052) Very commonly only 13 per cent is present, and

¹ F W Flot her states that a preparation made with sulphate of quimne contains less lime salts than when quimne hydrate is used, since the lime salts introduced in the water employed for reading the altaine form by what care retained by the latter, and are subsequently incupitated as calcium sulphate, unstead of remanuage in the feasibade product.

² To camera securate results, the sold solutions of the sample must be revised with a consultanable secure of amusuma, the volume of either or inhereform used aboutd equal that of the amusumand lapard, and the agention should be conducted unmentably, the netament with the advant should be repeated, and save must be taken that the whole of the prospetted shall due to simply for either. This occurs measurement by with any quinner, but if emobile the contraction of the contraction

⁸ The outgoind issue of the 1886 edition of the Drink Pharmacogonia required lips per ent of quinne, as estimated by the summons-etter process, but the entiscass of P. W Flether, O Umn ey, and others (Phorm Jou., [3], 233, 400) aboved that, if prepared according to the official directions, this proper team was impossible, and the amount was subsequently colored to 15 be cent.

4 Chas Umney (Pharm. Joss , [8], xvn 235) considers that, the BP. standard of quality being easily attainable, the manufacture of atrate of non-

occasionally (in the author's experience) from 9 to 11 per ceut, even in the case of preparations manufactured by English firms of fairly good repute. Foreign specimens sometimes contain only 4 or 5 per cent of alkaloid, and that not quinine.

The adulteration of citrate of iron and quinine is not limited to deficiency of total alkaloid, the quinine being sometimes replaced, without acknowledgment, by other emchona bases. The British Pharmaconoma prescribes no test for these, further than requiring the other-residue to be "almost entirely soluble in a little pure ether." The presence of these bases is best detected by dissolving the alkaloidal residue in sufficient dilute sulphuric acid to convert the bases into neutral sulphates,1 and treating the resultant solution as described on page 412 et sey To obtain reliable results a considerable quantity of the sample must be employed, but nearly the whole of the quinine is subsequently recovered as crystallised sulphate By separating this on a calico-filter, pressing it between folds of blotting-paper, and drying it at 100°, the anhydrous sulphate is obtained, and its weight multiplied by 1 18 represents the weight of the crystallised salt. If to this amount there is added 0 00133 gramme for each 1 c.c. of mother-liquor, a very fair direct determination of the quinine sulphate will be obtained, and by multiplying the result by 735 the corresponding amount of free quinine will be found.

In foreign specimens of citrate, substitution of the quinne by other emchona bases is common. Amorphous alkaloids are not unfrequently present in considerable proportion,

TINGUIND OF QUINING, BP, was formerly directed to be made by dissolving 160 grains of ergestilless clayhate of quiume in 20 fluid ounces of tineture of orange-peel, by the aid of a gentle heat, the solution being filtered after three days. This was an unasabstactary preparation, as in cold weather, or when too weak a punit was used, it was apt to deposit crystals of sulphates of quimne, and so after in strength. In some cave, at least, the deposit consisted largely of calcium sulphate. In the Phon macogenic of 1835 an equal weight of quimne hydrochloude is substantiated for the sulphate, so that the timeture is somewhat stronger than the old preparation. To determine the proportion of quinne in the timeture.

and quante containing only 13 per cent of alkaloid, nuless it alsos from some scordent, is a disgrase to pharmacy, and that any pharmacest who salls an attole of this character ought to be pumahed, unloss he can show good cause for the definency.

¹ Thus may be effected by adding a moderate excess of hot dilute acul, and then dilute ammonus, drop by drop, until the liquid is neutral to methylorange or limus. 1 fluid ounce should be concentrated, and shaken with ether to remove the essential oil of orange-peel. After removing the ether, the equeous liquid should be cooled, an excess of ammonia added, and then the whole shaken with ether in the usual way (see near 042).

Wrise or Quinsus, BP, contains I gram of expetallated sulphate of quinne and 14 gram of extra sead mosel fluid ounce of enange wine. It is a pit to be debased by partial omission of the quinne or its replacement by other cinchona alkaloids. For its assay, 2 fluid ounces may be concentrated to \$\frac{1}{2}\$ ounce, and then treated like the tincture of quinne (see above). If the skirold privalent modulum the step a mixture of chloroform and amplie all-bial of the best better More reliable results as only under the strategic deal of the step of the strategic deal of the step of the strategic deal and mostless of the strategic deal of

HYDROGUNINE, C_BH_LN_LO_s, was discovered by Hesse (*Eer* 'xv 856) in the mother-liquous from whein quame sulphate he energystallised, and subsequently in the commercial salt itself, in which it is sometimes present to the extent of a per cent. Quamic cannot be perfectly freed from hydroquamic can by repeated crystallisation of the neutral sulphates, but the syldicular quamic can be completely separated by converting the shikord into the acid sulphate and recrystallising this from wate or alcohol, when the hydroquamic remains in the mother-liquor

As preomitted from a cold solution of a salt by caustic soda, hydroquinine is amorphous, but gradually becomes crystalline. In the latter condition it contains 2 argus, which is driven off at 115°. From oblionform and other the alkaloid crystallises in dichests concentre groups of needles. It melts with darksming at 168°

Hydroquinine dissolves readily in alcohol, ether, chloroform, benzene and ammonia, but not in caustic alkali solutions, and is

only very spanngly soluble in water.

Hydroquinine resembles quinine in its lawo-rotation, fluorescence of its acid solutions, behaviour with the thallesquin test, and in its physiological action. It differs from quinine by only very slowly decolorising a solution of potassium permanganate.

Crystalline compounds of hydroquinine with cupreme, quini-

¹ The proportion of hydrogunuse in the lack is very small, and bears pure constant relation to that of the agumen. To obtain the hydrogunuse the alkalouis should be repeatedly cay-tallisted as send sulplates, the reaction quinnes get in do fly professions permangants, the hydrogunuse inheated from the filtered hydrogunuse in a state of the filtered hydrogunuse in th

dine, cinchonidine, and some other cinchona bases have been obtained, but not with cinchonine or hydrocinchonine

Hydroquinine has the usual well-marked basic characters of the curchona alkaloids $B_2H_2SO_4 + 6H_2O$ forms short prisms, soluble in 350 parts of cold water

The totrate crystallness with 2 agas in pussus which become anhydrous at 120° and are soluble in 545 parts of water at 17° C. The chromate is more soluble than the quinne salt, but crystallness with it, and can only be patially separated by bothing with water BEIG1+2 agas is reachly soluble. On mixing its solution with potassium include, the high-sociale separates as an only mass which gradually solubiae but does not become crystalline. The acid salt, $D(HI)_2+4$ aqua, crystallness in builhant yellow needley readily soluble in hot water to a colourless solution, from which the yellow salt separates again on cooling.

When heated to 140° with strong hydrochloric acid, hydroquining loses a methyl group, and is converted into hydrocu-

preine, Calla NoO.

When hydroquinine is bested to 140° with sulphuric acid containing 25 per cent of H₂SO₂ the alkaloud is unchanged, but when the dry sulphate is fused by heating it to 140°, the base is converted into amorphous hydroquinine in without alteration of weight or other change of composition.

Hydrogumene neutaises acids completely and forms some crystalisable saiss. When an ethereal solution of the base us gradually muxed with a solution of oxalic serd in ether, neutral hydroquintoine oxalite is formed as an amorphous brown mass, readily solution in chloroform, whereas the oxalate of quimense, obtained similally, forms a voluminous piecipitate, consisting of very minute needles

High equivars-sulphones and, Co₂H₁₆SO₄HiN₂O₃ + H₄O₃ no obtamed on dissolving phydroquames in cold concentrated stipliums soil. On diluting the solution with water and neutralizing it with ammoun, the sulphonic arcs separates in crystals, insoluble in other or chloroform and spaningly soluble in cold sola or aumonia in dulta scale it dissolves readily, forming crystallisable salts. The sulphurpe and solution is fluorescent and insponds to the challescount test.

Quinidine. Conquinue ConH24N2O2

This base is isomeric with quinine, and occurs frequently in cinchons barks (especially Cinchons Pitagensis) in association with quinine and other alkaloids. It also occurs in cuprea back, and is present to a considerable extent in commercial "quinoidine" Qumidine (see also page 393) crystallises from alcohol with 2½ aqua m large monochnic efforcescent pressure or needles From ether permanent rhomboledne contamung 2 aqua are obtamed, and from boiling water permanent plates with 1½ aqua. The whole of the water is duren off at 120°. At 160° the ambydrous alkalord begins to brown slightly, and melis at 168°.

Quandine sessables quinne in its isste and physological effects, in being despoted in hydridated crystals from alcohol, in it tolerably seady solubility in ethes, in giving the thillecopies of the solution in dilute subjeture seads and in the fluore-sense of its solution in dilute subjeture seid. It is distinguished from quinne by the permanent bulky preceptate its volutions yield on successive treatment with chlorus water, potassium fenicyande, and ammonn, and also by the very sparing solubility of its hydrivabile.

Quantiène Sulphate, B₄H₅O₄+2H₄O, crystelluses in white needles or long hard pussus when requise about 100 pasts of cold or 7 of boiling water for solution. It dissolves in 7 pasts of cold alcohol, and in 20 of chloroform, but is almost usoforble in ether, the cold alcohol, and in 20 of chloroform, but is almost usoforble in ether from the sulphates of the other cuchons. Alkholids in requiring a temperature of 120° to rendes it anhydrous, and in readily faking on the water sgum in most air.

Quinidine sulphate is an official remedy in the United States and France It is examined for other alkaloids by a test slightly modified from one devised by de Vill (Pharm Jour, [3], vill, 745), who utilises the fact that guindine hydrodide requires 1200 parts of water for solution. To test the purity of the commercial sulphate of quandine, 0.5 gramme is dissolved in 10 cc. of water at 60° C, and an equal weight of rodide of potassium free from any alkaline reaction added If the sample be pure, hydrodide of quinidine is procipitated on stilling and cooling as a heavy sandy powder, and if the liquid be allowed to stand for half an hour with frequent agitation and is then filtered, addition of one or two drops of ammonia will cause no turbidity in the clear filtrato A slight turbidity indicates a trifling admixture of other alkaloids, but if a decided precipitate occur the alkaline liquid should be shaken with a mixture of amylic alcohol and chloroform (see page 431), or chloroform only, and the solvent evaporated to ascortam the proportion and nature of the admixture, which may be cinchonidine or quinine. but is usually einchonine. The appearance of the precipitated hydrodude is sufficient indication of the presence of impurity, as in the presence of cinchonine or cinchonidine it is resinous instead of being sandy

For the detection of thorganic impurities (e.g., calcium or sodium compounds) in commercial quinidine sulphate, Hesse treats one gramme of the sample with 7 cc of a mixture of 2 volumes of chloroform with 1 of alcohol of 95 per cent Complete solution will take place in the absence of impurities

The presence of canhondone sulphate in the quindines all may be detected by treating the sample with pure chiloroform. Unless only a very small propersion of the impurity be present, part of it will remain unclassived. Smaller quantities may be detected by shaking the chiloroform solution with cold water, in which the whole of the canchondine and part only of the quantine salt will dissolve, and the formes will be prespitated on addition of Rochelle salt.

A solution of quantidue sulphate in chloroform is at first colourless, but on keeping becomes yellow with a slight green infliction

Ouinamine, C10H24N2O2

This alkaloid was first discovered by Hesse in the back of Cinchona succirubra, and has since been detected in C officinalis, resulenta, and several varieties of Crarhona Calisaya, particululy Ledon uma 1

Qumamme crystallises in delicate hair-like anhydrous needles, which melt at 172° C. Its rotatory power in alcoholic solution is + 104 5° for the sodium ray

Quinamine is nearly insoluble in cold water, more readily in boiling. Hot sloohol dissolves it freely. It also dissolves in boiling ether, petroleum spirit, and benzene

Quinamine theif is almost tasteless, but its solutions in consider avery printer. The solution in excess of dulute slightune said exhibits no fluorescence. And solutions of quinamine are very prious to decomposition with formation of an unouphous alliam of the solutions. It is solven that the property of the solutions of quinamine of the solutions of quinamine or produced, and undo costom conditions as pop un an in it, Ogli-Ha-NQ, results. When the sholl with childrene or bromme water and ammonia, solutions of quinamine yield a yellowesh amorphous precentate, but no green colour. The solut alkinoit, when moistened with strong interaction, gives a yellow coloration.

CONQUINAMINE, C19H21N2O2 occurs with quinamine, but in

¹ The motive-luques from the cystals of quants wilputs are pacquisted with Rechile six, the filtrate tractal with amonoma, and the preparation washed with other. The othered washings are treated with sectin said, and while wann treated with pleasure throughouts, till one cooling embourance on no longer be detected. Quanchas as then preceptated, tegeber with colouring metter. The filtered luqual to steaded with sods, and the reasons propputate dissolved in a nummum of hot 50 per cent, alcohol, from which quantum copatibles were cooling.

smaller proportion. It may be separated from the latter base by fractional crystallisation of the intracts, oxialities, or hybromides, the conquimamine salts being in each case the less soluble (Amalen, ceix 38, 62). Conquimamine forms colourless or golden-yellow tetragonal crystals, melting at 121¹-123², easily soluble in the chlowform, and benzone. S_p=+204 1º for a 4 per cent in alcohol B₂H₂SO₄+a squa is very soluble. The curvehloride is a yellow precipitate, becoming purple. Conquimamine closely resembles quinamine. When heated with concentified hydrochlorie each, it yields a poly un namine C₁H₂N₃N₄O².

Cinchonidine. C10H22N2O1 (See also page 392)

This base is contained in several species of circhions, but is especially characteristic of the red bark of *C succerubra*. According to D Hooper the absence of circhiondine is a distinctive character of Remina barks. It was formerly called quintdine.

Cinchondine crystallises in short anhydrous prisms on thin plates, soluble in 16 parts of alcohol and 188 of ether. It is reachly soluble in amylic alcohol and chloroform. It is here-rectatory, S_0 (where c=4 and $t=15^{\circ}$ C), in chloroformic solution being -70 C; while in dilute hydrochloro- and solution (c=5) $S_2=-174$ 6.

Cinchemdine resembles quinne in the direction of its optical activity, in the insolubility of the anhydrous neutral sulphate in chloroform, and in the sparing solubility of the tertrate in water. According to Hesse, it foins a crystalline compound with quinne containing C₃₀H₄N₃O₄ + 2O₃H₄N₃O_. It is distinguished from quinne by its lesser specific notation, its more sparing solubility in ether, its non-fluorescence, by not giving the thallesoquin reaction, and by the greater solubility of its neutral and acid sulphate and rodosulphate. The accurate separation of cinchendine from quinne presents great difficulties, and is discussed at length on page 411 et say. Cinchendine has only about one-fourth of the therapeuto activity of quinne

Cinchonndine is isomeric with cinchonine, from which it differs by its lawe-rotation, its greater solubility in other, the insolubility of its tartrate in water, the insolubility of the anhydrous sulphate in chloroform, and the formula of the crystallised sulphate

The normal saits of conchonadone are neutral to himus and methylorange, but acid to phenolphthalein. Thus the precipitated tartrate

¹ Omehonidime was formorly behaved to contain C_{th}H₁₈N₁O, but its convensor by heating with concentrated hydrochloria and into a post in ho nni-dine, C_{th}H₁₈N₂O, without formation of methyl chloride, and analyses of hydrochloride, sulphate, and ohloroplatinate establish the formula given in the text.

reacts to the last indicator like an equivalent amount of free tartance and, and the combined alkaloid can be estimated by titration in presence of alcohol with standard caustic soda or baryta Adhering Rochelle salt does not interfere

The following table shows the formulæ and solubilities of the principal salts of cinchonidine:—

			Solubility in Water	
Salt	Formula	Appearance	Cold	Hot
Hydrochloride,	BHCl+1 aq	Double pyramids	30	Readily
Hydrobromide,	BHBr+1 aq	Long colomiess	40	Freely
Sulphate,	BgHg804+v aq	Sliky lustrous needles, or thin quadratic prisms	100	soluble
Ovalate, .	$B_2B_2C_2O_4 + 6$ aq	Prismatic crystal	252 at 12°	
Tartiato, .	$B_2C_6H_6O_6+2$ aq	Crystaline pre	1205 at 16°	

Constantine sulphate, $B_c H_c S O_p$ is remarkable for the number of hydrates it is capable of forming. From a moderately concentrated equeous solution it erystallines with δ aga in building needles, from a host disconsisting of a gauge (official in the B and US Pharmacopenas), and from alcoloi in fine prisms or acclusing A hydrate containing δ agua (official in the B and US Pharmacopenas), and from alcoloi in fine prisms with Δ aqua A hydrate outsianing δ agua has been described by Hesse¹. The 6-stom hydrate is somewhat efficiescent, All water is lot at 100% and 2 agua re-absorbed in mosts for

Cinchondine sulphate a sometimes contaminated with an admixture of the contesponding salts of cutelouse and quundum To distect these, He s. s. (Zottade, Anal Chem, xx. 448) dissolves 0.5 gianume of the salt in 20 c. of vater at 60° C, and alids 1.5 gianume of Rochelle salt. a cystalline proceptate of the spreamyly solvible enchandment tatrate is produced. After standing one hour the liquid is filtered, and the fiftuate tested with a drop of ammonia. Any turnhelity or preceptate is due to the presence of more than 0.5 per cent of conclosure on 1.5 per cent of quantime These may be distinguished by beating the filtrate with potassium include as described on pages 413 and 426

Hager recommends the use of 01 guamme of curchondine

 $^{^2}$ Five commercial samples of our chonidno sulphate examined by A B Prescott, lost, at 100° C, proportions of water ranging from 6 36 to 7.01 per cent B₂H₃SO₄+3 aqua requires 7 30 per cent

sulphate, 03 of Rochelle sait, and 20 cc. of cold water. The leguid is frequently agistated, filtered after one hour, and tested with a few drops of ammonia. As thus performed, the test is less strict than that of Hesse, but perhaps, on that account, is better suited for medicinal purposes

The precapitate of caucionadine tartrate obtained in the above tests is soluble in about 1200 parts of cold water, but almost wholly insoluble in a strong solution of Rochelle salt. After drying at 100° C, it contains 80 84 per cent. of enchondine, It will contain quinner if any of that base were present in the sample. In such case the solution of the precipitate in excess of dulte sulphunic acid will be notably fluorespect.

Hese has also proposed to distinguish the stiphutes of the enchoins bases by their behavior with chloroform. The αnhydrous nentral sulphutes of quinno and enchounten are almost at each of the contract of

The presence of quantities and quantities in circle orders sulphate can be recognised by the thallesoquin reaction and the fluorescence of the solution in dilute sulphune and

Ησικοινισιουμπικ, $C_{10}H_{10}N_{10}$ (see also page 392), accompanes enchonables in many enchous batks, especially that of C rosulents, and passes into the dark sulphate mother-iquors in the quinne manifacture. It expressibles from alcohol in anhydrous piams, or from a chilate solution in leafests, almost insoluble in water, but soluble in chilatonia $B_{11}H_{20}O_{11} + C_{11}C_{12}$ crystallises from hot water in white needles, but from strong solutions the sail separates as a white mass, which after drugs greenholds magnesia.

If s s s states that homoemchonshino is an essentially different substance from emchondine, and that it is not possible to convert one into the other The two bases may be separated by fractional crystalization of their subplastes from aquous solution In presence of quinnes sulphate, the homoemchondine salt is said to crystalizes in the form of emchondine sulphate, the

Hydrocinchonding, C₁₉H₂₄N₂O, possibly identical with cinch onidine, occurs in the mother-liquous from homogenehousing.

Cinchonine C19HooN.O, or C9H7N C9H11(OH)N CH31

This important alkaloid is almost invariably present in cinclions basks. When the free bases are crystallised from alcohol the cunchonine is deposited before the quinne, unless the latter base is present in relatively large amount, in which case the greater part should be previously removed by crystallising the sulphates.

Cinchonine crystallises from alcohol in anlydrous shining prisms or needles. It melts at 165° C to a colourless liquid, and partially sublimes at a higher temperature. According to Hlasiwetz, it may be readily sublimed in a current of hydrogen or animonia

Cinchonine is almost insoluble in cold water, and requires 2500 parts of boiling water for solution.

One part of unchonine dissolves in 120 parts by weight of cold rectified spurit or 28 of boiling slockol, in 350 parts of chloroform, in 371 of other, and in 109 parts of amylta slockol, requires only about 13 parts of a mixture of 6 grammes of chlorofoniu with 1 of rectahed spurit, and is soluble in 23 parts of a mixture of 4 of chloroform and 1 of amylta elobol

A B Prescott found the following to be the solubility of cinchonine in different physical conditions, and at the boiling-point of the solvent —

	Parts by Woight of Washed Solvent requ			mod
Condition of Alkaloid	Réher	Ohlor oform	Amylio Alcohol	Banzone
Crystallined, .	719	828		.,
Amorphous, "Nescont," 2	568 526	178	22	276

It will be seen from these results that amylic alcohol is by far the best solvent for cinchonine, except a mixture of amylic alcohol and chloroform. On the other hand, ether is the best solvent for effecting an approximate separation of cinchonine from quinnine

When heated to a high temperature with an alkali, cinchonino yields quinoline, C₂H₁N (page 115), together with other products With sodime truchloride, cinchonine yields a yellow precipitate.

¹ The constitution of einchonine is discussed on page 168,

⁹ To obtain the alkaloid in the "nascent" state, the solvent was added to its sulphure acid solution, which was then warmed to the boiling-point of the former. The liquid was next made slightly alkaline with ammonia, shaken, kept warm for five minutes, and filtered

Cinchonine is not precipitated in the cold from a solution containing tartaire and by adding sodium hydrogen carbonate. On heating the liquid, however, carbonic and escapes and einchonine is separated.

The precipitate formed by ammonia in solutions of cinchonino is not soluble in excess of the reagent. The precipitate is amorphons when first produced, but speedily becomes crystalline

Condomne is sharply distinguished from quainte by the very limited solubility of the free base in ethe, by the solubility of the anhydrous insutral sulphate in chloroform, by its failure to give the fladleroquin reaction, by its dextro-rotatory power, and by the non-fladorescence of its solithon in excess of chiute sulphinic acid. Meshods of detection and separation based on these facts are given on pages 413 and 416

Čenchonsine Sulphates, (C.,H.,N.O.),H.SO.,+2H.O., forms short, hard, shunng, chin-chonther persans, with thothed summuts. The salt becomes anhydrous at 100°C, and molts with partial decomposition at about 240°C. Conchonium sulphate has a very bitter taste, dissolvers in 54 parts of cold water, and as reachly soliula (1 6) in alcohol. It is insoluble in either or benzene. The hardpraws salt is soluble in 60 parts of cold or 22 of boiling chloroform, a fact which distinguishes it from the sulphates of cinchonatine and unume.

A solution of cinchonne sulphate does not give the thallenouun reaction, and is not rendered fluorescent by dilution with very weak sulphuric acid

The mode of assaying of cinchonine sulphate is sufficiently indicated under the head of "Quinine Sulphate" (page 408 et seg)

Conclourse Hydrochlords, $C_0H_{\infty}N_c(0, HC) + 2H_0(0)$, is readily soluble in water and alcohol, and somewhat so in ether and chloroform. It has been not unfrequently employed to additioents sulphate of quinne. In such case the solution of the sample in very distate sulphane or nitrie and will give a white, circly procipitate of silver chlords on adding silver intrate. Cinchonine will be detected by the tests for that alkalout.

When heated in a dry test-tube, cinchonine hydrochloride gives purple fumes much resembling the vapour of reduce. The sulphates of the cinchona bases do not give this reaction

Hydrocinohonine, $C_{10}H_{24}N_2O$, is stated by H case to occur in cuprea bark

Chonorine, $C_{19}H_{26}N_2O$ (see page 392) is isomeric with cinchonamire (page 438). It dissolves very sparingly in ether (1 500). BHCl+2 aqua requires about 48, and $R_2H_2SO_4+12$ aqua about 35 parts of cold water for solution.

CINCHAMIDINE is a base probably isomeric with the above, and identical with hydrocinchonidine (page 430)

Amorphous Cinchona Bases.

Certain uncrystallisable alkaloids exist ready-formed in einchona barks, the proportion present being probably affected by sunlight and the piesence of any free acid in the bark

In the preparation of the salts of the alkaloids from cinchona bark, a further pointon of the bases undergoes conversion into a resincid substance known in commerce as "quincidine" or "amorphous quinine"

Gurschnes is obtained in quinne factories by precipitating the bown mother-hipous with ammona, and consists langely of two sikaloids, quinione and cinchononion, which are seemenwith and appear to be due to the school fleat on quinnie or quindine, and cinchonnes or cinchondriae, respectively. These amorphous products may also be obtained by heating the crystalhael bases in glycorn to a temporatus of 200°C, a 1cd substance being founded at the same time.

Commercial quinoidine is a dark brown, brittle, "extractiform" mass, softening below 100° C, and having usually a slight alkaline reaction. It is a product of indefinite composition which has never been very favourably regarded in this country, though it has received official recognition in the German and United States Pharmacopæias Both works hmit the ash to 0.7 per cent. By the latter it is described as almost insoluble in water, freely soluble in alcohol, chloroform, and dulute acads, and partly soluble in ether and benzene When triturated with boiling water, the liquid, after filtration, should be clear and colourless, and should remain so after addition of an alkali The German Pharmaconwia requires that quinoidine should dissolve clear in an equal weight of I part of dilute acetic acid with 9 parts of water, so as to leave scarcely any residue, and it must also form a clear solution with nine times its weight of cold dilute spirit Quinoidine is said to be liable to adulteration with mineral matters, resuls, liquonec, glucose, &c , all of which sophistications would be detected by one . or other of the above tests

For the purification of quinodine it is iscommended to digest the commercial article on the vater-lath, with 2 parts of benzene, while stirring or agitating. The clear solution is pound off, and the results washed with more benzene. The benzene solution is then shaken with a slight excess of dulte hydrochloine acid, the acid liquid separated, and rendered family alkaline by caustic soda. A portion of this solution is then tested for purify by dilution and sidition of a few drops of a concentrated solution of solution throughbut (hypoculphite), which ought not to produce any precupitate modulide on a further addition of water. Should impurity be undeated, the whole of the solution of quantities of hydrochloude must be treated with solution timesulphate as long as a permanent perceptate in produced. The buyul is then filled, warned, treated with excess of soda, and the precipitated quinordime washed with water and duried at 100°

Thus purified, quinodine appears in thin layers as a dast, yellowsh lown, transporent mass. It is completely soluble in bonzeno, alcohol and acids, and ether should dissolve at least 70 per cent of it. The normal sails of quinodine are said to have an alkalne reaction, and should be soluble in water in all proportions. When impure they form a clear solution in a httle water, but the lequid becomes turbal on further diduction.

To prepare a pure amorphous alkaloul, the eard sulphate of quanno or enchondune, according to the product required, is first rendered anhydrous by caucful drying at 100° G, and is then raised for a few minutes to a temperature of 130° to 135° G, when it melts and a wholly converted into the eard sulphate of the new alkalond

QUINICINE, ConHoaNoOo, is a yellowish, amorphous, anhydrous body, which melts at about 60° C., assuming a reddish-brown colour which becomes darker at 100°. It is nearly insoluble in water, but has a bitter taste. The alcoholic solution has a strong alkaline reaction, and absorbs carbon droxide from the air alkaloid is readily soluble in chlorotorm of ether Quinicine gives a green coloration when treated in solution with chloring- or bromme-water and ammonia, but is distinguished from quinine and quimidine by producing a white amorphous precipitate with sodium hypochlorite or solution of bleaching powder. In applying this test the liquid should be slightly, but not strongly, acidulated with hydrochloric acid. Quinicino may be separated from the accompanying alkaloids by adding ammonia, when the ammonium salt formed dissolves the liberated alkaloid, which may then be recovered by agitation with ether. If soda be employed instead of ammonia the alkaloid is thrown down as an oily mass

A solution of quinicine in excess of dilute sulphuric acid has a yellow colour but exhibits no fluorescence

Quantons forms crystallisable compounds with acuts, and double salts with the chlordes of planum and gold Neutral ozalate of quantons dissolves readily in hot chloroform, alcohol, or water in solution in a mixture of alcohol and chloroform the oxalate exhibits a night-handed rotation corresponding to a value of $S_2 = +26$ for the alkahol.

Quinicine solutions are not precipitated by Rochelle salt. They completely precipitated by adding excess of polassium thno-cyanate, which throws down quinicine thio cyanate as an oil which subsequently solutifies. It is soluble in pure woter, but insoluble in solutions of alkaline throeyanates

CISCOMMINIST, $G_{\rm q}H_{\rm mb}^{\rm N}$, $G_{\rm p}$ when precipitated by sola from the solution of no of its salls, forms a yallow viscous mass rought drawn out into colourless strings. It hquefices at about 50° $G_{\rm c}$ and at 80° times bown. At flaglo temperatures ($e_{\rm f}$, 100° $G_{\rm p}$) is becomes dark brown, and is converted into a substance resembling "quinoidine" U Topo cooling it remains $sG_{\rm f}$. As delixed from the rotatory power of the ozalste, in alcohole, aqueous, or chlore-forms colution, the value of $S_{\rm f}$ for enachromies is $\pm 90^{\circ}$!

In most reactions, including its behaviour with aumounceal salts and with hypochlorites, cinchomenne closely resembles quincine, and hence is distinguished from cinclionano and cinchonidate. It is distinguished from quincine by giving no gioen colour with chlorine- or bromne-water and ammouse

Cinchonicine is bitter, and in the free state has a strongly alkaline reaction. It neutralises acids perfectly, and many of the resultant salts are crystallisable

ANTIPRO-DASS Cetam anorphone bases, distinct from quincue and endomence, extra tendy-formed in eucliona barks. They are not convertible in guinetic or euclionence, and appear to be formed by the coalescence of two molecules of the opstabilisable alkaloids, accompanied in the case of quinne and quindine with the simmation of a molecule of water. Thus

$$\begin{array}{lll} 2C_{20}H_{24}N_2O_2-H_2O &=& C_{10}H_{40}N_4O_3.\\ &\text{Quintine or Quintiffine} &&\text{Diquimetine} \\ &2C_{10}H_{22}NO &=& C_{20}H_{44}N_4O_2.\\ &\text{Clinchouline or Christoper} &&\text{Distinchonleine} \end{array}$$

These bases constitute the greater part of the amorphous alkaloul contained in commercial quancibine. They are wholly amorphous, as also are all their saits. The solution of diquination in occess of diable sulpiture and is fluorescent, gives the thillengous reaction, and is dextro-rotatory. Disnucleonance does not possess these characters.

De $V(v_1)$ has pombed out a distinction between quintens, emchanisms, and the natural amorphous alkaloids. If the neutral exclusive of the bases be resultered milydrous by heating at 100° C, and the dry salts treated with chloroform, they behave in a characteristic name of Nathest of guaranteen dissolves spannight

in chloroform at the ordinasy temperature, but freely in the boiling liquid. On cooling, the solution deposits the greater part of the oxalate in crystalis. Anhydrous exhalte of cuchonicane dissolves freely in cold chloroform. By adding a few drops of water on the surface, the solution is transformed in a few nimutes into a solid mass. The oxalates of the natical amorphous alkaloids are very solublo in chloroform. The solution remains clear on adding a fow drops of water, but the water dissolves out some of the exalate from its chloroforme solution. The aniorphous exalate is highly deliqueseent, but the oxalates of quinterne and cinchomence remain unchanged in the air.

Alkaloids of Remina Barks.

The barks of the various species of Remijia vary greatly in the allicided which they contain. Thus, while the baik of R perlamentate contains qui nine and the allicid alkaloid our preine, that of R Perialization, which anatomically closely issembles the framen, and has been confounded with it, contains no alkaloid closely indiced to quimme except comparatively small proportions (0°1 to 0°2 per early of cinchonine and cinchonine and since of an annie. Cusconidine, which occurs in the bark of R Particiana, a lase found in that of Crachona Peliteira and, together with cusconine and aricine, which two bases do not appear to be present in Remijia baik. The bases moisted from this back by H asse west (in addition to cinchonine and candonamine) concussoring, chair-annie, concharamine, chair-annie, co

 1 To extract the whole of the alkaloids, amounting to 2 to 3 per cent . H $_{0.8}$ $_{9.8}$ treated the finely-ground back with hot alcohol, distilled off the solvent, treated the residue with excess of soda, and agreated with other. On shaking the separated othereal layer with dilute sulphure acid, a pale yellow, curily mass (A) separated, a portion of which remained suspended in the other and part in the yellow acid liquid (B) On senarating the latter (B) and adding very dilute nimic acid, emchonamine nitrate was precapitated (mixed with the minates of some of the bases of group A), while emchanic remained in solu-The curdy precipitate (A) was digested with dilute soils, the liberated alkaloids washed and an-dried, dissolved in hot alcohol, and treated with oneeighth of their weight of sulphune and (H.SO.) Almost all the concusponing immediately precipitated as sulphate, a small additional quantity separating on cooling Hydrochloride of chan amine was precipitated on adding strong hydrochlore and to the cold alcoholic mother-liquer. The filtrate from this was warmed and treated with a little potassium throcyanate, and the precipitate of concharramme throcyanate filtered off On adding more of the reagont, till the dark coloured solution became light brown, a mitchy mass esparated, after the semoval of which the solution was treated with excess of ammonia and shaken Concuscamidine does not appear to be a definite substance these alkaloids, like arieine and cusconine, contain four atoms of oxygen, and form a group only distantly related to emchanne and emchonamine. Concusconine has the same empirical formula as cusconine, aricine, and brucine, and resembles the strychnos alkaloids in some of its reactions. It crystallises with I aqua, and is dextro-rotatory, while cuscomine has a lower melting-point, crystallises with 4 agua, and rotates to the left. Concusconine resembles chairamine and its isomers in giving a deep groen coloration when the solution in hydrochloric or sulphuric seid is mixed with concentrated nitric acid, a reaction which is not common to cusconing or arreine Echstamine or ditaine, an alkaloid contained in the bark of Alstonia scholaris,1 only differs by Ha from chairsmine and its isomers, to which it presents a considerable resemblance Alstonine, Cat Han No Oa, an amorphous alkaloid, which occurs together with alstonidine and porphyrine in the back of Alstonia construta, is strongly fluorescent in acid solutions, and is not improbably related to the cuscondine group Hesse suggests that gelsemine, Co. Hos NoOs, the poisonous alkaloid from the root of Gelsemium semperonens (yellow jesamine), is related to these alkaloids, and points out that the coloration it gives with intric and somewhat resembles the reaction of concusconine

with benzene. The benzene was extracted with accts acid, and the acetic solution treated with a saturated solution of ammonium sulphate, which precriptated a mixture of the sulphates of characandine and conclaracandine, separable by factional crystallisation from hot water, in which the latter salt is the loss soluble.

DITAINS, or ZORITAININS, Qu'H₂NQ₂+4 aqua, foiras glossy prama Malta et 200°. S₂₀—28°. Very bitte Modenately soluble in water, alcohol, and ether. A strong lesse, not precapitated by ammonia Decompose sodium chiotois, setting fine causaire code. Rednesse Philing's solutions after bohing with hydronlicus acid. Onosentrated sulphunic seed dissolves ditaine with purple-wed colours; unitse and gives a pumple-wed, clauser; unitse and gives a pumple-wed, clauser.

DITAMINE, C16H15NO, an amorphous powder melting at 75°, soluble in alcohol, ether, and chloroform

RCHITENINE, $C_{20}H_{27}NO_4$, brownish, amorphous, melting above 120° Forms amorphous salts.

A full description of the alkaloids of Rennija Puriteena and Chenkhan Pelleturana barits has been published by O Hosse (Annalen, clxxxv 296, 323, ccxxv 211; Jona Chem Sec, xxxvn 165, xlvn 64, Pharm Jona, [3], xv 772. Arterin has been recently re-mresslegated by Médissan and Longrin (Compt Rend, cx 469) Ciuchonamune and cuprenne are described helow

CINGIONAMIRE, C₁₀H₂₁N₁O (see page 393), occus in the bark of Remajic Paradiavas (false capter but), a tree growing in Remajic Paradiavas (false capter but), a tree growing in the Columbian provinces of Anhoquie its selection is described on page 436. It is soluble in alcohol, ether, chloroform, benzene, and earbon disulphide, but only sparingly in water or petroletim spinit. It is very bitter, prosocously yields no methyl chloride when heated with strong hydrochloins ead; lyes no inestion with ferre chloride, and no colour with the thallecogum test. It is said to be insolible in strong hydrochlories call, but dissolves in strong nutries each with bright yellow, and in strong sulphura caed with the colour and the page of the province of the production o

Cuperris, $C_{\rm in}H_{\rm A}N_{\rm i}O_{\rm p}$ or $C_{\rm in}H_{\rm in}(O{\rm H})N_{\rm i}O{\rm if}$. This interest in glatical was discovered by F and and Go win loy in the bask of Ginchona cuprea or Remylia pedianellatia, a tree growing in the districts summoding the Manglelane Raves and the Upper Ginneco Since 1881, cuprea bank has been largely used for the manufacture of cumine ²

Cupteme crystallises from alcohol in the anhydrous form, but from ether in concentre pusses containing 3 agia. When the alcohole solution is district with vater, the precipitate contains B_g+ agia. The hydrates lose their vater at 123 $^\circ$ Cupreme is only springly soluble in ether or chloroform, but readily in alcohol. The alcohole solution is large-rotatory ($C_b=175~S_1$) alkining gives a dark reddish hown coloration with ferric chlorade, and responds to the thallenogum test. The solution of cupreme in diute sultrated by plume and is not fluoressent. The fice bees prequicted by

¹ Sée and Bockefontaine (Compt. Rend., c. 868) found cinchonamine (sulphate) eix times as toxic as quinne, cinchonidno, or cinchonime. An injection of 0.25 gramme killed a guinea pig in a few minutes.

³ For the preparation of express, the orasis quantum subplate from the copress that it dishorted in dishts solbitum seed, excess of custus coda added, and the quantum extracted by agriction with ether. The separated ribbins build as mentionless with subplance as viewed on the contract of the subplace as contracted with subplance seed, when express evaluation on column seed.

CUPREINE. 439

ammonia as only algebty soluble in excess, and may be extended by either When cupreme is liberated from a salt by a fixed caustic alkali, it dissolves on adding an excess of the respect, forming (with soda) a definite crystallisable composition denotating $(\nabla_{t}H_{t}^{N}X_{t})$ of Only, from the solution of which the alkaloid cannot be extracted by ether. This behaviour is due to the presence of a hydroxyl-group having a phenolic character (compare Morphine, page 311). The cup is in a test of potessium and solution are very soluble in water, and the corresponding compounds of calcium, lead, and silver have a strong alkaline reaction, and are most or less soluble in water. From the fact that alkahes from only mono-derivatives, while two atoms of the hydroxyl of explains can be replaced by scatyl, 31 is probable that the hydroxyl-stoms have different functions, as is the case with those of the morphism-molecule

When heated with hydrochloric acid (sp gr I 125) to 140°, cupreine is converted into apoquinine, without formation of methyl chloride.

The conversion of supression mod quitmen as described on page 398. Cuprators priefs two classes of saits Those of the general formula BÅ at a sparingly soluble, and the squeens solutions have a yellow colour, though their alcoholic solutions are perfectly colourless. The saits of the formula BÅ, are, as a rule, pretty freely soluble, and their sources solutions are colourless.

Oupseine Sulphate, E.H.SO,+6HLO, exptallases m mirate white needles, very difficulty soluble in cold water, and modulus in a saturated solution of sodrum sulphate $BH_2SO_1+H_2O_1$ exptallases in prams spannigly soluble in cold water. Oupseine three states from solutes efficiences in needle, very sparnigly soluble in cold water. Oupseine three-paraset is produced on adding potassium throcyanate to a hot solution of the monohydrochlorde. The liquid becomes turbuly and gradually deposite ascenial crystals of the salt. It is very sparnigly soluble in, and is precipitated in an only form by, an excess of the precipitant.

Homoquinine, When molecular proportions of quinine and

² When cuproun and causin potash or soda are mixed in molecular proper tions, a portion of the alkaloid (10 to 20 per cent) is extracted on agriction with other, but this may be prevented by using some excess of alkali.

^{*}DIAGRITI-CUPRININ, Oh.H.g.(C.H.P.)SN,Ob. was obtained by Hesse by heating curpense with secties subjective to 85° for a few borns. It forms heatsgonal plates melting at 85°, and as soluble in alcohol, chloroform, and other. The alcoholus colution is stongly slathnas, gives no colour with ferric chlorafe, but is turned dark green by chlorine and ammoons. By caustic allkalns, the base in hydrolyced in a few mustice with formation of equipment and accets and

cupreme are dissolved in dilate and, and the solution preoxpitated by ammonia and shaken with ethes, the solvent deposits on evaporation characteristic crystals 2 of a molecular compound of quantic and cupreme containing $C_{20}H_{24}N_{2}O_{2}C_{10}H_{23}N_{3}O_{2}+4$ agus. The same substance is readily obtamable by precupitating a solution of sodume cupremate with one of unume bridcohlorido.—

$$C_{u_0}H_{u_1}N_1O_{u_1}HCl + C_{t_0}H_{u_1}N_2O_1ONa \rightarrow NaCl + C_{t_0}H_{u_1}N_2O_{u_2}C_{l_0}H_{u_1}N_3OOH$$

This remarkable compound was disobvered and doesn'ted simultaneously by Howard and Hodgkin (Jour Olean Soc, Alon Olean Hodgkin (Jour Olean Soc, Alon Hodgkin Jour Olean Soc, Alon Hodgkin Jour Olean Soc, Alon Hodge Hod

The analytical differences between homogramme and corpreme have been fully described by Paul and Cownley (Pharm. Jour., [3], xv. 402).

Cinchona Barks.2

The back of various species of Chickens, which, with about thirty other alled genere, constitute the trib Chickense (Lee Rabiacos), have been long known for their antifolials proporties. These properties are chiefly due to pecture alladicate conclude therein, which alkaloids are alread through all the allied genera, except section species of Remire.

Nearly forty species of emchona have been descubed, many of which can only be discriminated with great difficulty. The emclosus form a very intractic genus, one series running into another through a series of intermediate forms, the number of which is limited to some extent in their native country by the fact that particular species are piactically confined to cortain distincts and obsentions.

Only some seven distinct species of cinchona yield bank of any practical importance. These are —

a Pale or Crown Barl, yielded by Cunchona officinalis (Peru) and albed species It occurs in quills, with a rough, blackish-brown or dark grey surface (For analyses, see page 446 et sec.)

² French; Ecoroes de Quinquinu. German; Chinarinden.

¹ Homogunine is deposited from ether in very thin pinematic lamine, having characteristically-shaped ends terminated with two oblique planes.

- b Yellow or Calisaya Bark is, with the exception of Ledger bark, the richest of all the cinchona barks. It now usually occurs in quills having a rough surface, but formerly was met with in flattened pieces known as "flat yellow bark"
- c Red Bark, from O rubra and O. succirula a, is distinguished by the ied colour of the sap and mature bark. It is extensively cultivated in India, and is remarkable for the large proposition of cinchonidine contained in it (For analyses, see page 446
- d Pitayo Bark, from C. Pitayensis, is imported in short, brownish, curly pieces, rich in quinin and quinidine
- e Columbian and Carthagena Bashs, from C lucumifolia and laneifolia, are imposted in soft quills of broken pieces of very variable quality Quinne is often wholly absent (Year-Book Pharm, 1888, page 426).
- f Ledger Bark, from Cinchona Ledgeriana, is the richest in quinine of all cinchona barks
- g Cuyven Bark, yalded by Remsjia peduarulada, as not a true onchona bark, and as the only known species of any other genus which yalds quinine, though the alhed alkaloid oinehon as mine (page 438) has been found in R Pas decama Cupren baik as peculiar m containing the interesting alkaloid ou prine 1 (page 438)

Hybrid banks are often produced, especially crosses between O officinalis and C succirulra (see page 447)

A concise description of the chief kinds of carchona bark, with their distinguishing characteristics, has been published by W Elboine (Pharm. Jon., [3], xiv. 653)

The British Pharmacopeua of 1885 gives the following as the characters of official (red) cinchona bark, from Cinchona successibra.—

"In quilla or more or less incurved pieces coated with the penderun, and varying in length from usually a few inches to a foot or more—the bark itself from about one-tenth to a quarter of an inch tink, or rarely more, outer surface more or less longh from longitudinal furrows and ridges, or thansvense cracks, amulting from longitudinal furrows and ridges, or thansvense cracks, amulting from longitudinal furrows and ridges, or thansvense cracks, amulting states, and blownish or redduk-hown in colour, numer surface brack-red or deep redduk-hown, irrigulaily and consider strated, fractum eneaty close in the smaller quills, but finally

¹ Formally, the anothona trees was murshably cut down and the back strapped off and durid in the sun or on huttles over a fire. A greatly improved plan is to make longitudinal measons in the lark of the growing tee, remove about half the bark, leaving the tennandes intact, and cover the stem with moss. First hint is then formed vary apidly, and this renewed back not only contains a larger percentage of total sikeloids than the original, but the alkaduse contains very much larger mojouton of quimne. fibrous in the larger ones, powder brownish or reddish-brown, no marked odour, taste butter and somewhat astringent "1

The characters which conventionally determine the marketvalue of "druggests' quills" are often vary fallacous, and have no relation to the real quality of the bark. A silvery conting on the engiderms of the bark is one of the points to which a factor importance is stacked, and renewed bark, though richer in alkalod than natural, does not sell readily for dregistric jurposes owing to the absence of the above characters, though it is readily bought by oursen manufactures.

A specimen supposed to be one of einchona bank can be readily identified as such by heating a small quantity in a test-tube, when a carmine-red or purple tar will be produced if the sample contain any of the anchona alkaloids

COMPOSITION OF CINCHONA BARKS

Cunchons balks contain, in addition to woody fibre, starch, guin, and mineral matters —the characteristic alkaloids; quinovin, and cinchons-red, cinchotsnine and quinic soids; colouring-matters, wax, fat, and knees of volatile oil

Water extracts only a portion of the alkalodal constnents of euchona bark, and a hot infason becomes turbul on cooling from the separation of sparnigly soluble enchotamates of the alkaloda. The solution obtained by treating enchosia tark with accidiated vater gives a white precapitate with tannin, a whitish precupitate with causies alkanes, and a yellow crystalline precapitate with platinic chlorida. Either of these precupiates yields the charcteristic colour of ou in oli in when subsected to div distillation.

The Ass of emchona barks from South American sources was found by Carl as to contain a sensible amount of coppe, but this motal was not detected by D. Hoop ar in the bark from frees cultivated in India (Pharm. Jour. [3], xm 545), though in other respects the general results are in agreement. The average total sist from mywarshs of 300 specimens of Indian bark was found by Hooper to exceed 3 per cent. Renewed and old natural barks contain less, but the proportion never fulls below 2 per cent. Towns and Ureach barks give as much as 4 per cent. of salt, and

¹ This description sefers to sed onchease bark in quilis, which, in the obtton of 1885, replaces the flat tel bark of South Americs, official in the Place resopura of 1897. The obtions judicocarly count to name the place of corner, whether South Americs, Malrias, or Ceyben, but they also count to recognize rail bark in sharings, although this as the form in which it was most commonly met with in commerce, and notwithstanding that the shavings are often much superior, as regard the onemoust of quanta, to other forms.

the leaves from 5 to 6 per cent From 24 to 27 per cent of the ash is soluble in water, and an additional 67 to 70 per cent in acid, leaving 5 to 6 per cent, of since insoluble

QUILOVIN, or CHINOVIN, is an indifferent body which appears to be a constant constituent of the cinchonas, but in a proportion solidom exceeding 2 per cent. It is dissolved on treating the bank with weak sods, and on adding hydrochloric acid to the solution is precepitated in admixture with quinovic acid and cinchonar-red. Treatment with milk of lime dissolves the quinovin and quinovic acid, which are representated by an each and separated by treatment with chloritorin, which dissolves the quinovin only?

Quinovin has recently been re-investigated by Liebermann and Gresel (Berichte, xvi 987, Pharm Jour, [3], xvi 987), who ascribe to it the formula CasHagO11 They believe two distinct modifications to exist, a -quinovin being present in cinchona bark and B - quinovin in cupies bark. a-quinovin is a white, very light, crystalline powder, quite insoluble in cold and almost insoluble in hot water, but soluble in cold caustic alkalies, lime and baryta water, and ammonia It is difficultly soluble in chloroform, other, and benzenc. It dissolves in nearly absolute alcohol (43 100 at 15°), and is obtained on evaporation over sulphuric acid as a gummy mass without any tendency to crystallisation, but it separates on diluting the solution with water in resettes of clear, very small needles. When precipitated by treating its solution in more dilute alcohol with water it is deposited in glittering white scales. The alcoholic solution of quinovin is dextro-rotatory (S = +566), does not reduce Felding's solution, and does not undergo fermentation with yeast. The powder is very bitter. In concentrated sulphuric acid it dissolves with orange-yellow colour and evolution of carbon monoxide Its solution in glacial acetic acid is faintly blue, as is also the precipitate thrown down on diluting the solution with

 β -quinovin closely resembles its isomer, but is not soluble in

[•] Quancum a prepared by Lue horm an a and Gussel form a preparednet obtained when the combons been are extracted from both by means of action Co. distributing off the alcohol, and treating the extract with a dilute numeral scal, the skizdowist are leadywed as alled The metalliche bowns reasons matter a digusted with warm milk of hims, and the filtered hynd presentated by hydrochebine and. The premptate is therefore a white powder. The bown leaves a hittle quanora scal undusorled as a white powder. The bown scalinous additional with water all a prespitate commonces to form, when small crystals of quinovus separate on standing. By resystallisation hours dulte alcohol tru to obtained pure in the form of small glittering scales.

absolute ather or sthyl acetate, and crystalines readily from chints clothd in handrome scales. In nearly absolute alsohol it insolves freely with slight avolution of hest, but after a time, even if evaporation be prevented, the greater part separation in glassy crystals containing $C_{\rm st}H_{\rm co}/1+{\rm EO}_{\rm st}H_{\rm co}$, which differeses very rapidly in the air with loss of the alcohol. The specific rotation of β -cumpon is +27 9°

When boiled for some time with dilute sulphures end, or, preferably, when their concentrated alcoholes oblitions are saturated with hydrochloric and gas and allowed to stand in a closed vessel for thirty hours, both the quinovins undergo complete decomposition into quinovins a cid and quinovit, a seschared body apparently containing Grand and quinovit, as sechared body apparently containing Grand closed to the session of the complete and the sechared to the complete and the sechared to the complete and the sechared to the containing the sechared to the containing the sechared to the containing the solution even after looling with acid. It is doubtful if quinovit has been obtained pure

Quivovic Acia, CasHasOa, is constantly present in cinchona barks in small proportion, and forms a snow-white powder of tasteless needles or scales, quite insoluble in water, ether, or chloroform, and only very sparingly soluble in boiling alcohol or glacial acetic acid. It is best dissolved by adding ammonia to the alcohol, and may be reprecipitated by acetic acid. Quinovic acid decomposes carbonates, and is soluble in ammonia and solutions of the caustic alkalies and alkaline earths, the solutions frothing like soap. The ammonium and calcium salts crystallise from alcohol in needles . the former salt losing its ammonia by exposure to air, or on boiling its solution. On adding an acid to an alkaline solution of quinovic soid, a hydrate of quinovic acid is thrown down as a very voluminous jelly, the whole contents of the vessel solidifying. In this form quinovic acid is very soluble in ether and alcohol. From the solution, the insoluble form of the acid separates in needles on standing. Quinovic acid gives with cupric sulphate first a green colour and then a precipitate, and the latter, when washed, has a bitter metallic taste When heated to about 300° C, quinovic acid yields pyroquinovic acid, carbon dioxide, and secondary products

 yields a solution of cuchotennie end. It is a yellow, amorphous, very hygroscopic substance, very soluble in watea, alcohol, and either, gives a green colori with feine chloride, is preceptated by starch, albumin, gelatin, and tarta-emetic, is hydrolyzed by ditule each since § 10 cos a of a cin ch on a-red, gives protectechuic and acctic acids on fusion with caustic potach, pieds: pylocatechic line of my distillation; and is readily decomposed in presence of excess of sikalies, with formation of cin china n-red The cunchotamates of the sikalonds existing naturally in cunchous bark are difficulty soluble in water, but dissolve readily in adultated water—pocked with ceremostate

Chromona-ngo or Chromoputuro Adup, Cu₂H₄,O₃. This is the natural colonum-matter of (red) annohous backs, from which it may be extracted by treatment with alkalnes. It are-piacipited from its red annuoused solution on addition of hydrochloric and The solution also yields a red precipitate with harmon chloride Cinchona-red is also produced by boiling cinchotanine and with chitts sulphure and, glinces being simultaneously formed. On fusing simultaneously formed. On fusing cinchona-red with potssh, protocated in a cit Q, CH₂O₈, is noduced. Cinchona-red is ussolution in water or ether, but sparingly soluble in alcohol. It is sometimes present in red bark to the extent of 10 per cent.

QUINIO ACID or KINIO ACID, $C_{\rm H_1}C_{\rm D}$ crystallises in well-defined hoxagound plates, fusing at 161^3 C. It has a strong and purely and laste, and is soluble in 2 purts of water, less soluble in alsoloi, and almost insoluble in either. Its solutions are isvortatory. When detilied with manageness thought and subphuze acid, kinic acid yields quinons, $C_{\rm H_1}C_{\rm D}$, which is deposited in deep yellow persons on the cooler part of the apprachas. This reaction was proposed by Stenhouse as a test for true cunchons bark.

The AKALODS are the most important constituents of cumbina backs, in which they exist in the form of cucho-banness and quantizes. The principal of them have sheady been fully described (page 398 et seg). The official interiors and liquid extent of emchance countries of the proton of the alkalouis of the back used for their preparation. (Pharm. Jour. [31], xv 445, 797, xv 463, 480)

Some kinds of cinchona back are occasionally wholly destitute of alkaloids Social specimens do not give a carnine-red tar when heated in a dry tube, this reaction being produced only when a cinchona base is heated with woody fibre.

The proportions of total alkaloids, as also the percentage of quinne, are extremely variable (see *Phan Jour*, [3], xiv 444, 445, 458, 797, 810, xv. 411, 453, 480), and chemical analysis

is the only means of forming an equinon as to the richness of a specimen of halk. De Vrij found the \$C. officinatis grown at Ootecammal to contain a proportion of total alkalouis varying from 11 96 per cent (of which 9'1 per cent was quinne) down to less than 1 per cent. Quinne so not seldom alsent from baries containing certain other of the cinchona alkalouis. This highest yield total alkalouis thrown is about 15 per cent. An Ootecamual bark has been found to contain 13\(\frac{1}{2}\) per cent, the greater purbang quinne, in eighty specimens of Colkieges Ledgerians, from Java, Moesa in 1879 found from 1250 to 109 per cent, of total alkalouis, the cunner manging from 11 6 to 0 9 per cent,

Of late years, owing to improved methods of cultivation, the proportion of quinne has sensibly increased. In the some species of cinchons, the natural bark, mossed bark, and renewed hask contain very different pascentages of quinner, the last being the richiest; beasies which the external conditions under which the treasgrown largely affect the relative and absolute proportions of the alkaloids in the bark.

Quinine and cinchonine are the emchona alkalouls of the most frequent occurrence Cinchonidine is bardly less common, and it occurs very largely in Indian led bark Quindline is not very frequent, and is never present in large amount

The following are analyses by D Howard of bank from cultivated cunchona trees grown near Bagota, United States of Columbia (New Granada). The characters of the banks have been described by E. M. Hollmes (Pham Jon.; [3], xxii 875)

Species of Cinchona	Quinnoe Sulphate	Qumme	Cinchout-	Quantine	Grachonine	Amorph-	Total Alkaloids
Thomsoniana,	5 94	4.45	0 27	0 20	0 82	0 74	0 54
Ledgerlava verdo,	5 90	8.08	0 00	0 20	0 01	0 41	4 43
Kogna,	7 90	6.48	0 00	trace	0 10	6 78	0 20
Moracka,	3 06	2.90	0 00	0 00	0 04	0 49	3 45
Tunn,	6 04	6.78	0 40	0 18	0 28	0 42	8 10
Pombiana,	5 88	4.41	0 34	trace	0 02	0 20	6 09
Officinalis,	6 82	4.74	1 22	0 07	0 10	0 42	0 50
Suctirchies,	5 93	4.16	2 77	6 02	0 12	0 20	7 72
Librid,	8 32	2.49	1 52	trace	0 64	0 52	4 07

¹ This is by no means a typical analysis of succirubra bark (see footnote, Page 427).

The following are analyses by D. Hooper, Government Quinologist, of cinchons barks grown in the Madras Government plantations, and shown at the Indo-Colonial Exhibition in 1886;—

Source of Bark		8 4 8	Smeltonidme.	Ime.	Smehonme.	Amorphous Alkaloids.	ng ago		
Species.		Description.	Quinme Sulphate.	Quantue.	Cimch	Qumqme.	Omob	Anor	Total
C succerubra,		Natural	2 57	1 01	1 14		2 11	0.88	6 04
,,		Mossod	2 27	1 09	1 68	1	2 08	0108	6 34
,,		Renewed	2 47	1 84	1 25	}	1 18	071	5 28
,, .		Branch	185	1 88	1 00		2 28	1 10	0 41
,, .		Root	1.66	1 24	1 43	0 41	0 77	1 27	5 12
,,		Renewed (shavings)	8 00	2 30	2 08		1 10	1 45	6 97
C robusta,1		Natural	1 02	1 43	1 58		2 08	0.31	5 40
,, .		Mossed	268	1 92	0 77		3 16	0.85	6 20
,, .		Renewed	5 92	4 40	0 51		2 54	1 65	9 10
		Branch	2 20	1.64	1 17	-	271	0.50	0 02
C micrantha,		Natural			102			0 40	2 32
33		Renewed	t1200	trace	1 12		2 45	1 02	4 54
,, .		Branch			1 00		1	0 45	2 05
C Calveaya,		Natural	1 62	1 21	2 18		2.32	0 20	5 05
»		Branch	0 79	0.50	1 93		0.78	0 48	8 73
C Anglica, 1 .		Natural	1 00	0.81	1 40	0 20	0.88	0 44	3 01
,, .		Branch	trace	trnee	201	0 25		0 38	2 05
C Ledgeriana,		Natural	7 88	5 49	0 82		1 33	0.98	8 52
19		Branch	2 07	2 21	1 07	1	0 49	0.50	6 27
O Javannoa, .		Natural			2 64	1:42		0 18	3 14
., .		Branch			1.40	1 48	}	0 45	3 87
Coffeenales, .		Natural	8 72	277	0.39	0 16	1.57	0 50	5 80
,, .		Morsed	4.57	8 40	0 15	0 20	1.50	0 02	617
		Renewed	5 00	4 21	0 65	0.22	0.85	0 70	0 03
galudiana,	11	Natural	0105	0 04	0 39		0 10	0.48	0 96
19		Renewed	0168	0 51	0 28		1 10	0.87	2.85
Pitayonels,	П	Natural	8 14	2 84	1 08	1 10	0.50	0.89	0 32
,, .		Mossed	5 12	8 81	1 91	0 08	0.95	0 87	7 67
,, .		Renewed	8 86	2 50	2 33	0 78	0 52	0 55	6 08
Humboltlana,	19	Natural	8 01	2 24	0 40	tance	1 55	0.90	5 18
,,		Renewed	1.72	1-28	0 48		0.04	1 07	8 42

¹ Cinchona robusta is a hybrid or cross between C successive and C efficuents, and C Aughes between C successive and C Calesque (W T This elion Dyor, Phasm Jour, [8], v. 481)

Analyses of a number of emehona barks from Madras have been published by B H Paul ($Planm\ Jou$, [3], xr 660) D H oper ($Pcar-Bool\ Pharma$, 1888, page 430) gives the following as the percentage proportions of alkaloids in typical barks from trees grown on the plantations of the Madras Government. \bot .

¹ In commenting on these results, B H. Paul strongly deprecated the preference given to the red bank over that of the crown and Calesaya bunks,

	Bark from C suptrub d.2	Crown Bark from C oftomake 2	Hybrid Barks
Quinine,	1 40	2.93	9 18
Cinchonidine, .	2 26	140	182
Quintôme,		0.08	0.04
Cinchonius,	1 92	0.42	1:17
Amouphous alkaloids,	0.68	0 42	0 50
Total, .	6.95	6 25	6 75

Hooper gives the following as the average centesimal composition of the alkaloids from numerous species of the above barks: 1---

	Rod Barks 2	Crown Barks	Rybrid Burks
Quinine, .	22.2	6510	41.2
Oinehonidine,	861	287,	40'9
Quimdine,		15	0.6
Cinchonine, .	50 9	80	97
Amerphous alkaloids, .	10.8	79	77
Total,	1000	100 0	100 0

which had sated prepotently on all some and. This prepulse had extended to the B. Thermanogouse, with the result that "every back preparation that appeared there was, in face, as officially indictated action," and continued for any papear discovery as in face, as officially indictated action, and continued for the comparatively valueless once (Fron-Fook Platris, 1888, pap. 449). The systal cores had, of which the analysis as given in the trip. Fall appears of the comparatively valueless once (Fron-Fook Platris, 1888, pap. 449). The systal cores had, of which the analysis as given in the trip. Fall appears of the core of the comparatively valueless on the systal core had a state of only underso quality, the propertion of adiabatic spiechess, and committed upon value burney from 3 to fige resident of subhation of possess, and considering less than 1 per cent of encohondriso. In the tool bark these proportions, were creased, the quame being usually 1 pap cent, with \$4, and 5 per cent of encohondrise. Bed bark bad becomes a drug in the markets, and almost worthless as a source of quame In registry of these encohones (Franz. Jon.) 3, xxx. 5049, Hooper pennted out that the fifty cave health of which the analyses, were grear were undeabloody of a typical chanatory, a back of the norther appears, as any such plants, and such administration on also chromatorized.

¹ See foregoing footnote.

² The mixed total alkaloids of red back have been introduced into commerce under the name of "Quinetam" This preparation is completely soluble in warm, strong alcohol, 81 grammes dissolved in 10 c.o of normal hydrochloric

ASSAY OF CINCHONA BARKS

The complete assay of the various species of cinchona buils, with the view of secretaining the proposition of the different alkaloids contained in them, is a process at once important and difficult. A great many mathods have been proposed, but very few can be trusted to yield accurate results when employed by chemists unused to them. Again, a process which is suitable when quinner unused to chief alkaloid presents becomes difficult of application when the canchomidne is in axoses. Unfortunately, also, certain processes which are extensively employed by professed quinologists are kept strictly scored:

In choosing a process of assaying cinchona bath, due consideration should be given to the kind of information required Thus, a pharmacest desuing to know the alkaloidal strength of ins buth will require a less eccuries and clabories process than a manufacture buying bark for the extraction of quinne. Again, in some case-th is sufficient to detenime the pracentage of total alkaloids, while in others it is very important to assortant the proportion of crystallised sulphate of quinne wholin the bath is capable of yielding. On this account, it is desirable to discuss the determination of the total alkaloids and of the actual comme seconstell.

a The British Pharmacoparia of 1885 presembes the following standard of quality and method of assaying 1 red emchona bark — "Test —When used for purposes other than that of obtaining

the alkalods or their salts, it should yield between 5 and 6 per cent of total alkaloids, of which not less than half shall consist of quinne and cinchonidme, as estimated by the following methods—

"1 For Quinne and Cinchondine —Mix 200 grains of red cinchona bark, in No 60 powder, with 60 grains of hydrate of

acid should give a clear solution, which, on addition of 2 grammes of Rochells shift must yield a precupitate squal in weight, after drying, to at least 65 per cent, of the quiestim taken —(From the Unofficial Foundary of this Dutch Society for the Advancement of Phanmacy, Pharm. Jone, [3], va. 662; "Quantum subject "occurs in common on a perfectly crystallised from

1 Based on a method devised by E R Squibb (Ephemeris, 1 106)

³This is not a very excelling requirement. Unfortunately no indication, as given of the proposition of sciental quinne which should be present. Consequently, one both may have double the instruser value of another, and shy to contain the required preportion of total alkaded, of which conclude the contain the required preportion of total alkaded, of which conclude shall consist of enchandrine and quinney, but shill only those of the less that falsoled to be present. As the shavings are excluded, and the established propulse as to the appearance of qualls tond to favore the use of instruct alkade that for notice network bart, the general effect is to promote the use of the least Volt. III. Park 19.

calcium, slightly moisten the powders with half an ounce of water, max the whole intimately in a small porcelain dish or mortar; allow the mixture to stand for an hour or two, when it will mesent the characters of a moist, dark brown powder, in which there should be no lumps or visible white particles Transfer this nowder to a six-ounce flask, add three fluid ounces of benzolated amylic alcohol,1 boil them together for about half an hour, decant and drain off the hourd on to a filter, leaving the powder in the flask, add more of the benzolated amyhe alcohol to the powder, and boil and decant as before, repeat this operation a third time, then turn the contents of the flask on to the filter, and wash by percolation with more of the benzolated amylic alcohol until the bark is exhausted If, during the boiling, a funnel be placed in the mouth of the flask, and another flask filled with cold water be placed in the funnel, this will form a convenient condenser which will prevent the loss of more than a small quantity of the boiling liquid Introduce the collected filtrate, while still warm, into a stoppered glass separator, add to it 20 minims of diluted hydrochloric acid, mixed with 2 fluid drachms of water, shake them well together, and when the acid hould has separated this may be drawn off, and the process repeated with distilled water slightly acidulated with hydrochloric acid, until the whole of the alkaloids have been removed. The acid liquid thus obtained will contain the alkaloids as hydrochlorates, with excess of hydrochloric acid. It is to be carefully and exactly neutralised with ammonia while warm, and then concentrated to the bulk of 3 fluid drachms. If now about 15 grains of tarterated soda, dissolved m twice its weight of water, be added to the neutral hydrochlorates, and the mixture stirred with a class rod, insoluble tartrates of quinine and cinchonidine will separate completely in about an hour, and these collected on a filter, washed, and dued, will contoun eight-tenths of their weight of the alkaloids, quinine and cinchonidine, which, divided by 2, represents the percentage of those alkaloids The other alkaloids will be left in the mother-liquor"

"2 For Total Alkaloids—To the mother-liquor from the proceding process add solution of ammonia in slight excess. Collect, wash, and dry the precipitate, which will contain the other alkaloids The weight of this precipitate, divided by 2 and

valuable kinds of bark for pharmacentral purposes. In the present Pharmaceneza definition, the quimme standard of canchona bark is reduced much below that of the 1867 edition, and only corresponds to a content of about 1 per cent of cumme

¹ Prepared by mixing 3 volumes of benzeno with 1 of smylic alcohol.

² It would be better to extract the alkalouds with obloroform.

added to the percentage weight of the quinine and cinchonidine, gives the percentage of total alkaloids"

b The following method of determining the total alkaloids of conchons bark is that of J E De Vrii, with certain modifications suggested by A B Prescott and J. Muter It is practically the official process of the United States Pharmacopera, and is applicable to all varieties of bark. Twenty grammes of the finelybowdered bark, weighed after drying at 100° C, is thoroughly mixed with 5 giammes of quick-lime and 50 cc of water. The mixture is then dried at a very gentle heat, not above 70° to 80° C When day, it is transferred to a flask fitted with an inverted condenser, and boiled with 200 cc of the strongest rectified spirit 1 The liquid is allowed to cool, and is then passed through a filter six inches in diameter, and the residue is again boiled with 100 cc of alcohol, and then washed twice with alcohol, using 50 c.c. each time. The filtrate is next rendered slightly acid by dilute sulphuric acid, and, after allowing any precipitate of calcium sulphate to subside, the liquid is passed through a very small filter, which is washed with a little alcohol The filtrate is evaporated or distilled till the alcohol is expelled, cooled, and again passed through a small filter, the precipitate, consisting of quinovic acid and fatty matter, being washed with water slightly acadulated with sulphuric acad. The filtrate, which contains the alkaloids in the form of acid sulphates, is then concentrated to about 50 cc or less, and transferred to a separator of 100 to 150 cc capacity. Soda is next added in decided excess, and the liquid containing the separated alkaloids then shaken without delay with 30 to 40 cc of previously washed chloroform After a few minutes' agretation, the liquid is left at rest till the chloroform has completely separated from the aqueous layer The lower stratum is then tapped off, and the watery liquid agitated three times more with chloroform, using from 25 to 30 cc on each occasion. The mixed chloroformic solutions are then distilled to a small bulk, the residual liquid evaporated to dryness, and the residue discil in the water-oven till constant in weight. The amount so found represents the total alkaloids in the 20 grammes of the bark taken. Cinchonine and cinchonidine readily become anhydrous at 100°, and quinine may be trusted to do the same Ounndine retains 2 agus in the water-oven, but the proportion in which this base occurs is too

¹ The spirit may be methylated, but should be proviously dehydrated to about 98 per cent by being kept in contact with fieshly-ignited potassism carbonate. A Soxhiet's tube or equivalent arrangement might probably be advantageously employed for the alcoholo treatment described in the text.

small to affect appropally the accuracy of the assumption that the alkalouds are weighed in the anhydrous state. If preferred, however, the temperature may be raised to 115° C 1

For the assay of yellow cinchona bank, ether may be substituted for the chloroform employed in the above process

c The following method of assay is due to Prollius (Archiv d. Phasm, cax 85, 572), with precautions suggested by De Vrij, Biel, and others It is practically the process of the German Pharmacopæia (1882) —Prepare a mixture of 85 parts of other (sp. gr 0724 to 0728), 10 parts of alcohol (sp gr. 0830 to 0.834), and 5 parts of ammonia (sp gr 0.960), all by weight, making 100 parts in all Treat 10 or 20 grammes (according to its supposed richness) of the previously dried and very finelypourdered conchons bark in a tared glass-stoppered bottle with twenty times its weight of the above solvent-mixture, observe the exact weight of the bottle and its contents, and agitate at intervals during four hours If any loss of weight occurs, add sufficient of the solvent-mixture to restore it, agitate and weigh again Carefully decant into a flask as much of the solution as can be poured off perfectly clear, and ascertain the quantity taken by re-weighing the stoppered bottle Distil off the ether, evaporate the residual liquid in a tared beaker at 100°, and weigh the residue when thoroughly dry Then -

Weight of solvent mixture employed × weight of residue Weight of alkaloids solution decented fin bark taken

The crude alkaloids thus obtained are dissolved in thitse hydrochloric sord, the solution filtered, and the filtrate made alkaline with caustic soda and repeatedly agriated with chloroform, which is separated, exponenced, and the residual alkaloids weighted after drying at 100° in the usual way De Vij found the junified alkaloids so obtained from a red Java baik to be 83 5 per cent of the total cude alkaloids previously extracted

With a few modifications of muon importance, the method described in the sext is that used by most quantity of the bark (about 2 lis). Having itsels to work on a very large quantity of the bark (about 2 lis). Having itsels to work on a very large quantity of the bark (about 2 lis). Having itsels, with lims, sleohol, and soul in the mannel described in the tast, he to equintize the aqueous solution of the entiphates with sole, filtens, washes sightly, dissolves the prosystate in actic and, and filters from any uninsolved colouring-matter. The filtrate is divided into two equal parts, A and B A as meruphated by ammons, filtratel, and the litrate sisken with B A as meruphated by ammons, filtratel, and the litrate sisken with The work of the litrate is the litrate sisken with the colouringth of the litrate is the litrate in the litrate sisken with the litrate is the litrate in the litrate in the litrate is the litrate in the litrate in the litrate in the litrate is the litrate in the litrate in the litrate in the litrate is the litrate in the litrate in the litrate in the litrate is the litrate in the litrate in the litrate in the litrate in the litrate is the litrate in t

d The following method for the estimation of the total alkaloids of cinchona bark is that of Hager The accuracy of the method has been confirmed by O Medin (Zeit Anal Chem, vni 477, 447).—Ten grammes of the dried and finely-powdered bark are treated for a short time with 100 cc of water and 10 grammes of caustic potash solution of 1 35 specific gravity. The mixture is then heated and kept at the boiling-point for a quarter of an hour. Fifteen grammes' weight of diluted sulphuric acid (sp gr 1 115) is next added, and the whole boiled for twenty minutes. After cooling, both hound and residue are transferred to a measuring cylinder, and diluted with water till the whole has a volume of 110 cc1 The liquid is then passed through a dry filter, and 60 cc. of the filtrate (=6 grammes of bark), mixed with 50 cc of a cold, saturated, aqueous solution of pierce acid. After standing for half an hour the precipitated picrates are filtered off, washed with a little cold water, dried at 100°, and weighed ' The product contains 42 5 per cent of its weight of alkaloids, calculated as quinine A preferable plan is to suspend the washed precipitate in cold water, add excess of caustic soda, and agitate with chloroform. The chloroformic solution of the alkaloids is then treated as in process b. The picie acid method of assaying cinchona barks is said to be accurate, easy, and expeditious

Separation of Cinchona Bases.

The separation of the alkaloids of einchona and allied barks is an extremely complex operation, and as respects the sarer alkaloids outside the scope of this work. But the accurate separation even of the commoner alkaloids, such as is frequently required for commercial purposes, is very difficult, and its accurate performance presents special obstacles to an inexperienced analyst. In some cases it is sufficient to determine the proportion of civitallisable quinine, which may be effected as described below, but in other cases it is necessary to determine also the emchonine, cinchonidine, and occasionally the quinidine, quinamine, and amorphous alkaloids. For the separation of quinine from the admixed alkaloids, ether is usually employed, but it must be remembered that the separation effected by this solvent is not an absolute one, all the free cinchona bases being more or less soluble in ether, especially in the presence of quinine The anhydrous sulphates of quinine and cinchomdine are almost insoluble in chloroform free from alcohol (see page 430). but in presence of sulphate of cinchonine or quinidine sensible

 $^{\rm 1}$ This is allowing 100 c c. for the liquid, and 10 c c for the bulk of the residual woody fibre, &c

quantities pass into solution Crystallisation of the quinine sulphate from water affords a simple and fairly accurate mode of separation, which has the advantage of being similar to the proeess employed by the manufacturer, and hence is regarded by many as furnishing the best proof of the yield likely to be obtained in The following method of separating the quinine in the form of sulphate is described by J Muter (Analyst, v 223) -Treat the total alkaloids, or the ether-residue from 20 grammes of bank, with warm distilled water slightly acidulated with dilute sulphuric acid, till the mixture is perceptibly acid. Add water to make 70 c c for each 1 gramme of alkaloids taken, and then very drivet sods with constant staring tall the hound is exactly neutral. with a faint tendency to acidity Digest the liquid at 85° C for five minutes , then cool, and leave at 15° C for one hour Filter the liquid through a small double filter (24 mones diameter), the two filters being previously trimmed to equal weight, and receive the filtrate in a graduated cylinder. Wash carefully with water at 15° C till the filtrate and washings measure 90 cc for each I gramme of the maxed alkaloids. The filter and contents are now completely dued at 100° C, and weighed, the second filter being used as a counterpoise. To the weight in grammes add 000817 gramme for each e c of filtrate and washings. The sum divided by 0 855 gives the corresponding amount of crystallised sulphate, and this number multiplied by 5 gives the crystallised quimme sulphate obtainable from 100 grammes of dried back

The quaine sulphate so obtained is agit to contain emblandino sulphate, and should be tested for this admixture as directed on page 412. The remaining alkstoids may be recovered from the mother-liquous by concentrating the liquid somewhat, adding solar excess, and agitating with cholorform. On exponenting the olloroform, the bases will be obtained in a solid state, and may be separated as described on page 459.

¹ The solubility of the cundiona bases in ether at 15°C is given by A B Prescott as being—for quinine, 1 25, quindine, 1 30, emchandine, 1 188, and for cunchomic, 1 371 The amorphous cunchone alkaloids are teadily soluble in ether.

	ı	ž
	-	44.0
		å
	1	
	1	4
	1	3
	1	19
		1 01
		post
	1	mde
	1	2
3	1	Snel
	1	2
	1	fate
	1	900
	1	000
	1	200
3	1	of a
4		-
	1	- Contract
ì	1	9
ı	1	3
1	1	9
ì	1	-
	1	1
		7
	J	1
		-
		7
	1	
		4
	1	
		400
	1	1

0	B The Richersal Bolution is errip every disophed in 10 parts of pro- tion an alcoholic solution of not troes of coline must be cretelial herepathic is munciately profit herepathic is need as ease only a registrance. In such a case only a well sturred, and left twelve hours.	A wealt of set less that and preferably E grammes of the mored althouse no fees take a may providency, and breakd in a closed stell while the form the breakd is a closed stell while the form at the fillering, and the restitue while the waited with a small quality of elect.	To reaction the property, and provided, and the almost there will be a provided to the control of the provided and the control of the property of the provided and the control of the provided to the provided and	Properties A. Commerce of the control challenge and Park. To control challenge and the challenge and th	A weakle for the law from a haddy weakle would be considered by the second considered considered considered by the second considered consi
---	---	---	--	--	--

The precapitation of the quantum as herepathitie is stated by D a vid H or var I to give securate issuits in aktiful hands, but, instead of throwing down the quantum from a sulphune and solution by functure of nodine, De Viri recommends, in his more recent papers, the use as a precipitant of the noisoliphate of the aniorphous cinchona bases commercially known as "quinodine". This forms a readily soluble toolousliphate, and by employing a previously nepared solution of it any error from the formation of periodised prodesultwate of quantum is avoided! De Viri directs

1 Pharm Jour, [3], vs 461, xu 601 One part of commercial "quinosdine" is heated on a water-bath with 2 parts of benzone, whoreby the quinordine is partly dissolved. The cold, clear benzene solution is shaken with excess of dilute sulphune acid, an aqueous solution of the acid sulphate of quinoidine being thus obtained. The amount of alkaloid is then determined in a small portion of this solution, and the lest is slowly treated with I next of rodine and 2 of potassium rodide dissolved in water for every 2 parts of amorphous alkaloid known to be present. The rodine solution must be added very gradually, with vigorous stirring, so that no part of the quinoidine solution shall come in contact with excess of iodine A flocculent, orangecoloured precipitate of iodosulphate of quinoidine is formed, which by shight elevation of temperature cuagulates to a dark brownish-red resinced body The yellowish liquid is poured off, and the precipitate heated to 100° with water, when the hand is poured away The adhoring moisture is evaporated off at 100° C , when the iodosulphate remains as a soft and tonacious mass, which becomes brittle on cooling One part of this substance is dissolved by heating with 6 parts of alcohol of 92 to 95 per cent. The solution is allowed to cool, filtered, evaporated to dryness, and the residue dissolved in 5 parts of cold alcohol When filtered, the solution thus obtained is ready to: nee

In using this solution for the determination of crystallisable quitine in a mixture of cinchona bases (as free as possible from einchonidine), 1 but by weight of the alkaloid is dissolved in 20 parts of alcohol of 92 to 95 per cent , containing 1 5 per cent of sulphune acid (H2SO4), which amount is sufficient to convert the bases into acid sulphates. The solution is then diluted with 50 parts of unacidulated alcohol. To this liquid, at the ordinary temperature, the rodosulphate of quinoidine is added drop by drop from a burotte, with constant stirring, as long as a dark brownish-red precapitate of heropathite is formed. As soon as all the quinine has been precipitated, and a slight excess of the reagent has been added, the liquor acquires an intenso yellow colour The beaker is now covered and heated on a water bath till the liquid begins to boil, and all the procipitate is dissolved, when the liquid is allowed to cool After standing twelve hours, the beaker is weighed with its contents The hand is next passed through a small filter, leaving the crystals in the beaker, which is then again weighed to ascertain the weight of the liquid. The crystals on the filter are washed back into the beaker, and as much alcohol added as is necessary to dissolve the crystals at the boiling-point When quite cold the beaker is again weighed, the recrystallised herepathite collocted on a small filter, and the empty beaker again the addition of the resgent to the solution of the mixed alkaloud of cunciona bark, but it has been pointed out by Chils tons en. Shimoyama, and others (Pharm Jow, 13), xm. 441, 1016, xvi 205, xvi. 554), that cunchomidine, if present in notable quantity, is lable to be precipitated along with the quinne, and hence this base should be separated as completely as possible by a previous either-ineatment, as directed on page 455. The use of the iode-sulphate of quinodine prevents any subsequent isolation of the

amorphous alkaloids of the bark under examination.

Instead of converting the quinine in the ethereal solution B into herapathite, David Howard (Watts' Dict Chem. 2nd ed. 11 177) agritates the ethoreal liquid with excess of dilute sulphuric acid, and, after heating to boiling, adds dilute ammonia till neutral to himus, using no more water than is necessary On cooling, the quining ciystallises out almost entirely as sulphate, which salt is almost insoluble in a cold solution containing ammonium sulphate The crystals are filtered off, washed with a little cold water, pressed between blotting-paper, and dried at 100° C 734 parts of the anhydrous salt represent 100 parts of the crystallised sulphate The product should be tested for emchaniding (page 412), which may be present in small quantity. The alkaloids existing in the mother-houor from the quinine sulphate are then recovered by concentrating the liquid somewhat, adding soda in excess, and shaking with chloroform. The bases are extracted from the separated chloroform by driute acetic acid, and the solution treated as in A

The mixed alkaloids of yellow cinchona bark consist chiefly of quimme, and hence the portion soluble in ether represents the most useful constituents of the bark. Pale and red barks, on the other hand, contain a considerable proportion of alkaloids insoluble

weighed. The difference indicates the weight of the mother liquor, which is added to that of the main quantity

The recrystalhead batepathne obtained as above a wealed on the filter with a stimical solution of herepathne in alcohol of 92 per cont. The adhening liquid is removed as far as possible by pressing the folded filter and its concents between blotting-paper, and the filter in them an-dried. The preceptates is then detached from the filter, due at 100° till constant, and wagind. The amount found is corrected by the adultion of that remaining in solution, as accretional by orderisation from the weight of the mother-inquer One hundred greatmes of alcohol of 92 per cent dissolve 138 gramme of heavpathnet at 24° C J, and 128 grammes at 16° O

The weight of herepathite found, multiplied by 66066 gives the anhydroue, or by 07409 the corresponding weight of crystallised, sulphate of quinno Instead of drying the occayabilised herepathite, it might pooledly be three with standard sodium throsulphate solution. 21.58 parts of sodium thus found

represent 100 parts of herepathite.

or sparingly soluble in ether. Hence the use of chloroform in the general process for assaying circhona backs (see page 451).

In some cases, the alkaloids soluble in ether are contaminated to a considerable extent with colouring matter. In this event, the following is a good method of obtaining colourless aumine sulphate -The ether-residue is dried thoroughly and woulded It is then dissolved in 30 c c of absolute alcohol, and decinoimal sulphuric acid cautiously added from a burette, till the liquid is neutral or very faintly acid to htmus-paper or methyl-orange Each c.c. is comvalent to 0.324 gramme of anhydrous alkaloids The bound is next evaporated nearly to divises, and a measure of doesnormal subhuric acid added equal to that previously required for neutralisation Thirty c.c of hot water are added, and the haund houled till complete solution acsults. Purified animal chargoal is next added, in quantity equal to the weight of the ether-residue, the hauid heated on the water-bath for twenty minutes, filtered, and the residue washed twice with boiling water acidulated with sulphuric acid. The filtrate is brought to a concontration of 70 cc for each 1 gramme of ether-residue taken, and thon cautiously neutralised with caustic sods, and further treated as described on page 451

Instead of commencing the separation of the alkaloids by other, Moon's recommend; that the neutral solution of the nursul alkaloids should be treated with excess of solution of potassimi and matching the (Rochelle sail), which throws down the quinne and enchonidine as tertakes. The same procedure is adopted in the British Pharmacopian (see page 460). The proepitated interface are washed with a hittle cold water, decomposed by excess of alkali, and the quinne and unchonductne separated by ether, it equinne dissolved being either discolding the processing of the pro

The estmaton of the relative proposions of quinno and cinchomidine in the mixed tarinates, by observing the optical activity (page 410), has been recommonded by several chemists, but in practice it is difficult to obtain the alkaloids in a sufficiently jure condition to render the results trustworthy.

The following method for the separation of the cinchona bases insoluble, or enerly insoluble, in either may be applied to the residue left on teatment of the mixed alk-louds with either, as in De Virly's process (page 464). It may also be applied directly to the mixed alkinolad extincted from a sample of bark, in which case it may be carried on simultaneously with Muter's process for the production of crystallined quanne sulphate as described on page 454.

Separation of Cinchonine, Cinchonidene, and Quinidine ANALYSIS OF CINCHONA BASES.

We make the contracted from the bank or from the filtest from the organized quantum suphate by treatment with toch another contract and the remains and the contract of the remains the contract and the remains and the contract and weaking the contract and the contract and the contract and the contract and weaking being collected in a gradient quantum. The mixed-alkaloids extracted from

ҹ The Precipitate is dried at 100° to 105° C and weighed, the outer filter bong used as a counterpose. The amount found is counterfold by adding 50088 gramme for each 1 c o measured by the filtrate and wash-water. The

has not previously been oparated, the amount of crystallised sulphare found must be multiplied by 245, and the product subtracted from the weight sum multiplied by 0.797 gives the weight of carellonidine II quinne of the tartrate before calculating it to A preferable plan 19 to with dilute sulphure and, add ammons, and extract with ether, weighting or titrating the alkaloid (See also dissolve the precipitate off cmchondine

• The Fitness was consistent of the foreignal bulk, cooled, a drop of diline action and alded, and then cases of a summarized the first two and parameters build from the constitution was consistent and an alleged from the constitution of the first foreign and the first of the given rod are produced [now] though the present the first two first foreign of the given rod are produced [now] the first two first foreign on a double confine point of the first and it is

д

thed carebonnes is extracted by agration with choosing, which is expressed, respectively and the residue wagind on thereto. The for such 1 oc measured by filteria A, and 00006 for each c o of hiteria B. Any Filtrate is measured and made distinctly alkaline with caustic soda, and the precipiamorphous alkalond may be dissolved out by spurt of 0 94 specific gravity The Properties is deed at 100° B and weighted it weight is contowed by the addition of 00077
and weighted its confined by the addition of 00077
and weighted its confined by the same mailtributing (B). The same mailtributing (B) and the presentation may be decomposed the presentation may alliated at the presentation in the alliated extracted by eight, and

Weighed age also,)

uthor's method described on next page.

459

The foregoing process, with experience, gives very good results, the sum of the separated alkalouls frequently amountaint of 90 per cent of the mixed bases operated on It is well suited for the seasy of Indian barks. The least satisfactory part of the process is the separation of the enchonine from the amorphous bases has dilitie spurt. A cautious employment of ether would perhaps be preferable. If the process of separation be conducted simultaneously with the determination of the crystallised quinne sulphate in another proton (page 404), the whole analysis can be combileted in about six hours.

According to II selbing (Fharm. Zeatesh f. Russland, 1888, Analyst, XII 907) the presence of much quantum prevents the complete prespitation of the emehandine and quinne as tartintes, while the prespitate with polassium noide, if tenesious or resultant sunstand of crystalline, contains unadomie, with or without quintend (It seems more pubblish that the resumes precipitate constate of the hydrochides of amorphous silkaloids, which can be kept in solution by modests addition of alcohol 1

The directions in the foregoing table can be modified with considenable saving of time by tistning the alkalouls and their satis instead of weighing them. Thus, for the determination of the concloundars, he weaking of the precipitate durints with cold water should be omitted, and the filter containing the precipitate and the adhering Rochelle salt solution immersed in boling wides. A drop of phonolphthalem solution is then added, and the hypothetic distribution of the containing the containing the precipital position of the containing the

An axacly smular method is applicable to the treatment of the preemptate produced by potassaum rounds. This should be washed with a little of the preemptant metaod of with water, and then immersed together with the filter in boiling water. On titrating with \(\frac{T}{2} \) alkala and phenolphicalise used it co of the standard solution required represents 0 0162 gramme of quisidizes precripated as hydrodide.\(\frac{T}{2} \)

The chloroforme solution of the canchonian may be directly tirated with standard and and methyl-orange (see p 131) mstead of being evaporated to dryness, but, of course, the amount found will meliide any amouphous alkalond also extracted by the chloroform.

² This procedure does not dispense with the necessity of making a correction for the amount of quindine lost in the mother-liquor and washings

BERBERINE AND ITS ASSOCIATES.

Berbeline is an alkaloid occurring in a very large number of plants, in many cases in association with one or more of the alkaloids, berbamme, oxyacanthine, hydiastine, canadine, &c. It is the only natural basic colouring matter receiving practical amplication as a dvs.

The principal sources of berberine and the associated alkaloids are the roots of the following plants —

PLANT	Alkaloms, &c
Berberts vulgarus (Barberry),1	Berberine, ovyacanthine, berbamine, and at least two other alkaloids (Hesse)
Berberus aquafolium,	Benberins, 2 %5 per cent , oxyacauthine, 2 %5 per cent
Coptus trujolus,	Berberine, 4 per cent
Coptus testa (India),	Berberine, 8; per cent , coptinine (crystallis able, G10ss)
Hydrastis Canadensis (Golden seal),	Berbermo, 1 3 to 1 8 per cent, hydrastine, 1 5 per cent, canadine, xanthopuccine, &c. Also meconin and phytostestin
Jatenhuz Calumba or Cocculus palmatus (Unlumba 1001),	Berborine, columbic acid, and the neutral principle columbin.
Mensepermum Canadense,	Berberine, oxyscanthine, menispermine, men ispine

Berberine has also been found in Wood unipa, a yellow dyewood fiem Uppen Assam, in St John's wood, from Rio Grands, in Berbers aristata, Caulophijlism thalatrivaks, Coseinium fraseivatum (Ceylon Calumba wood), Calcolene polycarpa, Podophijlism pellatum, Xamthorhasa aquifolfa, and Xanthozpism dassum Hacules II yd rashine cours also in Sylphophorum diphishum

Berberine, ConH17NO4, or C18H11(O CH2),NO.

Berbeime is solated from the root of Hydraclis Comadensis by boiling with water, evaporating the decection to an extinct, and exhausting with strong alcohol One-fourth of its volume of water is added to the filtered alcoholes solution, the alcohol distilled off, and the residue treated with dilute sulphure acid Berberne sulphate crystallises out, and is decomposed by freshly-precip-

¹ A concentrated liquid extract of barberry root still recorves a limited application for dyeing silk and leather yellow. In America, the root bark is commonly used, but in Europe the entire root is generally employed.

tated hydroxude of lead. The alkaloid may also be converted into the sparingly soluble intrate or hydrochloride instead of the sulphate

L. Wolff recommends a previous treatment of the root with petroleum ether to remove fixed oil

Berberne may be solated from barberny or calumia nort by exhansing the material with selocil, evaporating off the sprint, taking up the resulue with wates, and treating the fillested solution with crosses of hybrichlorise ead, when between hydrochloride crystallises out. The salt may be purified by resolution in alcohol and rescontiation by either?

Berberne czystalniese with difficulty in small, concenticulty grouped prisms, or bright yellow, silky needles? When an-drosi, the crystals appear to contain 5½ agun (W II Perkin, jun), of which 3 agus is draven off at 100° At this staepersture the crystals lose their baste and become yellowish-hown, at 110° the clauge is very rapid, and above 160° total decomposition cocus. Flot iman n gives 120° as the melting-point of berbeine, but Porkin considers this figure too low 8°

When warmed, berberine emits a faint but peculiar odour resembling minone

Berberme has a persistent, very bitter taste, and is employed medicinally in doses of 2 to 5 giams. Sixty grains have been taken by man without mjury, but the alkaloid is poisonous to dogs and other of the lower animals.

Berbenne dissolves in 500 parts of cold water, and more readily con boiling. The solution is neutia to litmus II has spannight solutile in cold, but readily in bot alcohol, and in anylic elocid. Berbenne is slightly solutile in chloroform and benzene, and insolute in either (separation from oxyscanthine and hydrastine) and petioleum spart. It is said to be taken up with difficulty

¹ Berbenne may also be prepased by preceptating an squeeze decoction of barbary root with lead sociate, and treating the concentrated filtrate with excess of milphuro and. The preceptate of behavior sulphate is washed with cold water, and separated from lead sulphate by solution in boiling water, which on cooling deposit the sail in yallow needle.

² An orango colour, or other shade darkes than length yellow, is indicative of impurity

⁴ E Sch mid has obtained some evidence that berbeine piequied from the commercial sulphate is occasionally a mutime of borbeine with methylberbeine. He obtained pure between becomes ting the alkalon the sections compound, B_CH_CO, from which the free base was liberated behating in alcohole solution. This obtained, between contained 6 que, all of which was leat at 100° C. The anhydrous alkalod scutouly began to darken below 150°.

from its acidulated solutions by amylic alcohol, chloroform, and benzene 1

When treated with a fixed caustic alkal, berboime is colouned brown, and on boiling a resinous mass separates On distilling berberne with milk of lime, quinoline is formed Fusion with caustic potash produces borberic acid, C₈H₈O₉, and an acid of the composition C₈H₈O₉.

When boiled with excess of furing hydrodic acid, two methyl groups are aluminated and a sait of berberoline, $C_{\rm B}\Pi_{12}({\rm OH})_{\rm B}NO_{\rm p}$ formed On rendering the duluted liquid slightly alkaline by ammonia, an intense blackash-blue coloration is obtained, probably owing to oxidation. Nitro acid gives, with herberoline, a magnificent violet coloration, which on standing or warming changes to a deep redtable-livowi

Concentrated nitra coid dissolves berberne to a daix, reddishbrown liquid, which on dilution with water gives a yellow flocation precipitate partly soluble in ammona. If the daix solution of is bebrein in strong mains and be warmed condition mixedly convey, with formation of be 1 be roin a acid (a pyridine-inarrhovytic soid) rage 113, bx x1; a call, and other products

Potassuum penmanganate in presence of potassuum caibonate oxidises berbeinne with formation of hemilplinic acid, $C_{10}H_{10}O_{0}$, and other products (W. H. Porkin, jun, Jour Chem Soc, J. v. 71)

By the action of nascent hydrogen, berberne is reduced to hydroberberine, ConHo, NO.

Herberne duselves in concentated sulphure and with orangeyellow colon, changing to they geen on warming On adding potassium bachromate, or other oxubising agent, a black colour is changing to robel (or brown-robet changing to brownish-yellow) is obtained. Froble's reagent gives a brown or green colour with baberne, or, according to Hirse sh hau see, in a mimediate yellow, changing through dails brown to violet-lawn. Sulphovanadas acid is stated to give a fine wide tooloution.

According to E. Schmidt (Plans Med., 1887, page 649), bebours he as temakable tendinery to combron with norths oliveria, such as alcohol, the nectors, and chloroform, to form crystaline compensate. When bestern and chloroform are mived in molecular proportions, they units to form a beautiful crystalline body, parmanent at 100°. This does not appear to be a beautiful crystalline body, parmanent at 100°. This does not appear to be a beautiful crystalline body, parmanent at 100°. This does not appear to be been and chloroform, such as the act decomposed by noisi study into beherne and alcohoroform, such as the such decomposition profits of the latter Berteins and also combine with a second moleculo of chloroform, but thus best his water of crystallivation. Schmidt has also described a compound of betterine with acctors, of the formula Call-NO, Call. Of

Berbernne is also characterised by the insolubility of many of its salts (eg, the chromate, picrate, hydrodide, chloroplatinate, autochloride), and the sparing solubility of others in presence of excess of mineral and

On pouring chlorine-water (avoiding excess) on to a solution of berberine strongly and dated with hydrochloric or sulphuine and, a zone of bright ad colour is formed at the junction of the liquids, and is still recognisable as a pink coloration in a dilution of 250,000 i

On cautiously adding todused potassium rodde (avoiding excess) to a solution of a berberne salt, BHI₈ is thrown down as an extensely misolible red-brown precriptate, which caystallises from strong alcohol in red needles, or on adding water in green irridescent scales which completely polaries light

Mayer's reagent yields with herbenine solutions a precipitate of the approximate composition B_2H_2 HgI₂, containing, after drying at 100°, from 50 to 52 per cent of the alkaloid

SALTS OF BEDERRINE

Berberine is a weak base, but forms definite and leadily crystallisable salts with acids. The salts have a bitter taste, and are mostly very sparingly soluble, the pyrophosphate and acetate being exceptions

Berberne Nitrate, BHNO, separates in fine yellow needles on accidating a warm aqueous or alcohole solution of berberne with nitre and It is soluble in about 500 pairs of cold water, more readily in hot, and almost insoluble in alcohol or water strongly solutiated with nitric and It does not darken or undergo other change at 100° km.

Be be use Hydrochlorade, B.HCl+2 ag, as prequitated in golden yellow needles on adding hydrochlora ead to a warm angus solution of the alkaloid. It requires about 500 parts of cold water for solution, and is almost unsoluble in alcohol or distal hydrochlorac ead. The salt is with difficulty decomposed by bases, the bleasted alkaloid burn gap to retain chloran Peolonged dipestion with ithangs fails to decompose it completely, lut sulver cords caudily decomposes the solution. Telebrem hydrochloride derivation to an onange colour when heated to about 60° C, but regains to coping and colour on cooling. By prolonged exposure at 10° in colour changes permanently, and much of the salt becomes reachly solution in the colour solution and the colour.

BHAuCl4 is amorphous, brown, and quite insoluble in water

¹ Brucine gives a similar reaction with chlorine-water, but the original solution is colourless, and the reaction produced less permanent than with betterine It crystallises from boiling dilute sloohol in chestrut brown needles, unchanged at 100° B,H_FtCl₀ forms a yellowish precipitate, sinost insoluble in all the ordinary solvents. It may be crystallised from boiling amylic alcohol, in which it is slightly solubla.

Be berine Hydrodude, B.HI, obtained by precipitation, forms minute yellow needles, extremely insoluble in cold water or possisium indice solution. It does not darken or suffer other change at 100° BHI₃ is precipitated on causiously adultan todassed potasseum indice (carefully avoiding excess) to a solution of a berberne sait in hot spint. It is quite insoluble in cold water When recrystallised from hot alcohol, the smaller crystals transmit light which is completely polarised (compare Herepathite, page 403)

Berbe ins Sulphate, B.H.S.O., is met with in commerce both in the morphous state and cystallised. The latter form, which is considerably the higher pixed, can be prepared by dissolving 15 grammes of the amorphous preparation in a boiling mixture of 250 c c of alcohol with 8 of acetic serd, when on coching the crystallised salt separates out. It has an oange colour, and is pointment in the air when free from impurity. It is soluble in about 100 just of water According to J U Li o q d (Ame Drug, Supt 1884), the yallow crystalline powder obtained by heating commercial beberne sulphate with ammonia and shalting with either is not the free alkaloid, as commonly assumed, but a neutral sulphate, B₂H₂SO_o which is readily collible in mater.

B,H₂CrO₄ is obtained in orange-yellow needles on adding potassium bichromate to a boiling and very dilute solution of a salt of berberne. The salt separates entirely on cooling, and is extremely insoluble in cold water or an excess of the precipitant

Berberine Pienate requires 45,000 parts of cold water for solution. As a consequence, on mixing aqueous solutions of berberine and piene acid in equivalent proportions and filtering, a liquid is obtained free from yellow colour of bitter tasts

Betherme Acetate is prepared by adding beherme sulphate to a solution of a potassum acetate in textified spirit, and heating goatly till the yallow salt has dissolved. After cooling, the liquid is filtered from the potassum sulphate, evaposated to a syruly, and shaken with either, when berberine acetate, B(C,H,O,by, is projected as a crystalline orange powder. It is readily solution water and alcohol, nearly insoluble in either, and loses acrid on exposure to an

OXYACANTHINE, C18H19NO₈ This base is contained in Berberie vulgaries, and remains in the mother-liquor, from which the berberine vol. 111 Part II. 2 G

has been separated as hydrochloride. The liquid is treated with caustic soda, when a dark-coloured precipitate is thrown down, from which ether dissolves oxyacanthine, berbamine, and an unnamed alkaloid, while another brown-coloured amorphous base remains undissolved. The ethereal solution is treated with acetic acid, and the resultant acetate decomposed by sodium sulphate, when oxyacanthme sulphate is precipitated, berbanine remaining in solution On decomposing the solution of oxyacanthine substate with ammonia the free alkaloid is precipitated in flocks, which after drying at 100°, melt at 138°-150°, but when crystallised from alcohol or other it forms anhydrous needles which melt at 208°-214° Oxyacauthine is readily soluble in chloroform and benzene, but only sparingly in petroleum spirit. It may be separated from berberne by extracting the ammoniscal solution with ether or chloroform. From its acidulated solutions it is not extracted by petroleum spirit or benzene, and only spaningly by chloroform Oxyacanthine is dextro-rotatory in chloroformic solution (for 4 per cent at 15°, S, = +131 6°) BHCl+2HaO forms small colourless needles, the 2 per cent aqueous solution of which shows Sn = +163 6° Hot strong solutions are coloured green by ferrie chloride Oxyacanthine closely resembles narcotine. Like morphine, it reduces indic acid. Concentrated sulphuric acid, with or without molybdie acid, is stated to give no colour at first, but on standing or heating a vellow colour is developed, according to L v Hirschhausen (Zeit Anal Chem, xxiv 163), Fiohde's reagent gives an immediate violet coloration, changing to yellowish green at the edges

When heated with caustic potasi and a little water, expeciation mells to a brown mess which floats on the fused cilical. This consists of the potassium derivative of $\beta - \infty y$, a can it him elements of water. A similar change occurs very reality even at the ordinary temperature, by the action of alcoholic pota-th or buryts on $\alpha - \infty$ -yascanthine. Rither fails to extract the β -modification from the alkalme solution. Hydrochloire and prespitates $\beta - \omega y$ -capacitine, which is soluble both in alkalics and excess of acid. With much acid, $\alpha - \infty$ -yascanthine hydrochloire is precentated.

Bereaming, C_{IS}U₁₀NO₂₀ the second Berbe is alkaland soluble mether, was obtained by He as a Gerichte, xix 3190) by adding sodium intain to the liquid from which oxyacenthine had been thrown down as sulphate. The precipitated bedamine nitrate when decomposed by ammonia yields a crystallino precipitate of the field should be a supplied to the field should be a supplied to the field should be a supplied to the state of the state

tanning 2 aq and melting at 156° The salts are crystallisable and readily soluble $B_2H_2PtCl_0$ is yellow, crystalline, and only slightly soluble in water

Hydrastine. $C_{21}H_{21}NO_6$, or $C_{19}H_{16}(O~CH_3)_2NO_2$ (see also page 470)

This interesting base occurs with berbeame (and canadine) in the root of Hydrastis Canadensis or Golden Seal? Forms found 1½ per cent in the direct root, but the yield in manufacture is from ½ to ½ per cent. It also occurs in Sitylophorum diphyllium. Hydrastis from berbeamen in being coloriles, but commercial medicand preparations of baberine from Hydrastis are not unfrecuently called hydrastine.

Hydrastane forms colourless or milk-white four-sided pusms, melting at 132° and decomposing at a ligher temperature with an odour of phenol

Free bydrastine is tasteless and oloculess, but the salts have an acrd teste The alkalont is the cheft if not the only active principle of Hydrastis Canadensis 1 Its posenous in lange doses, 3 grains being fatal to a frog in four immutes. It resembles strychime in causing death by aniesting the respiratory movements in a tomic spasm

Hydrastine is insoluble in water, and nearly insoluble in alkalino solutions. It dissolves in 120 parts of alcohol, in 13 parts of chloro-

2 F. W. il le Im evitacis the coarsely-provided root of Hyphotata Connocionary with boiling water annihilated with scone and, exportes the decotors to a gyrup, and adde excess of duite mlyhurus and Atte standing, the berberne sulphate whule cytablises out is filtered off, and the filtrate scentiales of with ammous. The prespitates contains much hydrastine, and on again filtering and adding excess of ammount to the filtrate structure prespitate a predicted, which is suid to contain connocions. Both preceptates when hoteled with ethylenetic growth and the suid to contain connocions. Both preceptates when hoteled with ethylenetic growth and the suid to contain connocions. Both preceptates when hoteled with ethylenetic growth and the suid to contain connocions. Both preceptates when hoteled with ethylenetic growth and the properties of the suit of

Ever hard t purifies hybristime by dissolving the freshly-pocupated akkadi an animum of bohing chloreforn, filtering though gisse-wool, and pouring the solution into excess of cold alcohol. On shaking the liquid vigicously for some minutes, the hydrastim experiates as a fine crystalline precipitate, which is subjected to a repetition of the process and resrystallised from boiling selected.

² The root of Golden Seal is a bitten tonic analogous to calumba. It is exhibited in the form of powder and in doses of 8 to 24 grains. The hydrochionides of the mixed alkaloids of golden seal are sometimes sold under the name of "hyd a stine".

form, in 16 of benzene, and in 83 of ether. It is quite insoluble in petroleum spirit. The solubility of hydrastine in ether may be utilised to separate it from berbeine. Hydrastine is levo-rotatory, S_A in chloroformic solution (1 2759 gramme in 50c c) being — 57 8⁻¹.

Hydrastane is a feeble base, and is completely extracted by chloroform from solutions freely acidulated with hydrochloric acid In part, however, it is dissolved as hydrochloride, which salt is very soluble in chloroform

With the exception of the picrate, the salts of hydrastine are generally uncrystallisable, or are obtainable in crystals by special means only Most of them, except the tannate and piciate, are soluble in water, the solutions having an and reaction

Hydrestine hydrochloride and sulphate are used in medicine. B.HCI is best prepared by passing dry hydrochloric and gas over the surface of a solution of hydrestine in sulpricos and alcoholfree other. After drying over sulphure and the picepitate brine a micro-crystalline powder essily soluble in water and chloriform a micro-crystalline powder essily soluble in water and chloriform. B.HSO, is smalarly obtained by continuity adding a solution of strong sulphure and in either to an othereal solution of hydrestine. The salt readily takes up water and forms a gummy mass.

Hydrastine solutions give no colour-reaction with chlorine-water, With nodused potassium rodide they yield a deep brown flocculent precipitate

Hydrastine may be approximately determined by titration with Mayor's reagent (page 141), but the precipitating power of the solution is materially affected by the dilution of the liquid

Pictic acid forms in hydrastine solutions a yellow amorphous precipitate of the picrate, BA+4 aq, which is deposited in splendid yellow needles from its solution in boiling alcohol

Solutions of hydrastme are precipitated by potassium bichromatic on touching the separated precipitate with a diop of strong sulphunc acid, it instantly becomes bright red, the colour fading in a few seconds This behaviour distinguishes hydrastme from strychnine and gelesomme (nege 36).

If a solution of hydrastane be sodulated with sulphune acid, and a few drops of a docunormal solution of potassuum permanganate added, the colour of the reagent is instantly discharged, and an intense blue fluorescence is developed. A single drop of a 1 per

¹ The figure for specific rotation given in the text is that of Figure 1 and Will. Eukman practically confirms this. F. B. Power (Pharm Jour., [3], xv 298) gives the widely different figure —170°

² The crystalized sulphate of hydrastane adventaged by some manufacturers as sumply sulphate of benbeune, to which the name hydrastine is persistently missipplied.

cent, solution of hydrastine when treated in this way renders a large test-tube of highed strongly fluorescent (A B Lyons, Plan , Jour, [3], xxx. 880) Excess of permanganate must be avoided, or both the alkaloid and fluorescent product will be destroyed. The fluorescent body differs from sesculin in not being extracted from either acid or alkaline solutions by chloroform or either and in not having the fluorescence intensified by addition of alkali¹.

The colour-reactions of solid hydrastine have been re-investigated by A. B. Ly on a (Plazm Jon. [3], zv. 880) with the following results —Concentrated sulphuric acid dissolves the pure alkalout with faint yellow colour, changing to a deep blue-puiple on heating. If the respent contains a trace of inthic acid a yellow colour is produced, and with a large proportion (1 1000) the colour is criage-red. Pure intin acid produces a permanent orange solution, which on adding water deports an invaluble substance, and yields a liquid exhibiting an intense blue fluorescence (compare last pare).

With sulphure and and oxidising agents (compare jage 368) with inangeness discribe an energy delicities of colour-reactions. With mangeness discribe an energy delicities of the street and the charges, and beaming through camme to a yellowah shod of red, which after a time changes inther suddenly to a pale orange-yellow. This reaction distinguishes hydrastine from strychimon and gaisemine, while berberne dissolves in sulphure and with yellow colour, changing on addition of the oxidising agent to violet, then to choosite-brown, and finally becoming orange-red (then intermediate chocolate-brown stage distinguishes the berbernie exection from that given by strychimon). Potassium permangenate gives with hydrastine and sulphure and the same colorations as manganesed touck, but the changes are more mpid. A violet time is sometimes produced of the the red is developed, the contrary order beam characteristic of strychime.

Fronde's reagent gives with hydrastine a sage-green colour, slowly changing to brownish, and then gradually fading. This succession of tints is very characteristic. Sulphovanadic acid gives a rosered colour, which fades slowly

On treating an acid solution of hydrastine with exidising agents (eg, manganese dioxide and sulphinic acid), it splits up into opianic acid (page 298) and hydrastinine, a base closely resembling cotamine (page 299). If the exidation be effected in

¹ The same fluorescent excitation-product is sometimes developed in solutions of hydrastine by mere exposure to air. No suther pure hydrastine nor any readyformed constituent of Hydrastis root appears to be fluorescent;

alkaline solution, the action proceeds further, the chief products being he mipinic (page 299) and niotinic acids (page 111) This behaviour suggests a closs relationship between hydratine and narcotine, but hitherto all attempts to convent one of these bases into the other have been unsuccessful.

Hydrastinine, C,H,1NO,+H,0, produced together with opining acid by the action of oxidising agents on hydrastine, forms white crystals, melting at 116°-117° C, or at 100° after heating for some time to that temperature. It dissolves in water to form a strongly alkaline and very bitter solution. It is also soluble in ether, ethyl acetate, benzene, and petroleum spirit, and crystallisus from each of these solvents with I aqua, which, however, is elummated in the salts, a fact probably due to the formation of a closed 1mg. C1H,1NO,HCl crystallises in feebly coloured needles, soluble in water and alcohol The aqueous solution is optically mactive and feebly fluorescent B, HeSO4 forms yellow crystals showing a green fluorescence, and is soluble in alcohol. Hydrastinine, when treated with acueous potash, yields hydrohydrastinine, CnH12NO, and oxyhydrastinine, C. H., NO. The latter is a feeble base, melting at 97°- 98° and distilling above 350°, and forms crystallisable salts. The former base is also formed by the action of reducing agents on hydras-It forms white crystals melting at 66°, and yields civstallisable salts

When hydrastanne is caulised in dulte alkalma solution with a solul attracted abutton of polosseum penamagnate, it is converted almost quantitatively min oxyhydiastinine, $C_{11}H_{11}NO_3$. Excess of the oxidiama agent and slight heating carries the oxidiation to hydrastinine area $(L_{11}H_{11}NO_3$ and body crystallising in flat needles melting at 164° , soluble in alcohol and ether, and yielding to preceptate with silver, bainum or lead salts 3

Canadine, $O_{21}H_{21}NO_{\phi}$ is an alkaloid accompanying benerine and hydrastine in golden scal root. Until recently these was some doubt as to its actual existence, L1 o yd having failed to detect it in the extract from a very large quantity of the root, but F

¹ E. S. h m of t consides that naconine contains three melberry groups and hydrastene only two, their formule beng respectively Q₂H₁₂(Mol₂)NO₁ and Q₂H₂₁(Mol₂)NO₂. As these bases both yield on conduction opinine and, which contains the methodry groups, and colarance contains the and group, it follows that hydrastine contains the methodry discounting the contains of the analysis of the constitution of a methylated hydrastime.

² The constitution of hydrastimine and hydrastine has been the subject of varieus investigations by Freund, Will, Ross, Resemberg, Leolman, Schmidt, Wilhelm, Keistein, Heim, Philips, Dormeyer, and others (Berachts, xix. 2797). Wilhelm and E Schmidt have independently isolated the alkaloid, which is described as forming five, white, shining crystals, melting at 134°, and more readily soluble than berberine in water and alkahes

The salts, with the exception of the sulphate, are soluble with difficulty in water and alcohol. B, HCl and B, H,SO₄ are crystalline By treatment with odine in alcoholic solution, canadine is converted into the hydricodide of methyl-berlevine, and hence it probably has the constitution of a d thy d ro in cft p lene berb evine

XANTHOPUCCINE is the name proposed by Lerchen (1878) for an alkaloud of doubtful existence occurring in hydrastis Liberators in snowbile in ether and chloroform, but soluble in alcohol and hot water. The alcohole solution yields light brown stangles with toolsed potassum nodde

Indications of other alkaloids in hydrastis have been obtained by A. K. Hale (Year-Book Phanm, 1874, page 31) and J. C. Burt (ibid, 1886, page 95)

Calumba, or COLUMBA, 1 is the root of Jateurki.a Calumba or Cocculus palmatus, a herbaceous climbing plant occurring in the forests of East Africa

The calumba of commence consists of dried transverse slices of the root. It possesses mill bither teme properties, and the tineture, extract, and infusion are official preparations. The roots of buyonia and Frasea Walters have been occasionally sold as calumba.

and Frasera Walters have been occasionally sold as calcula Calumba root consums three distinct bitter principles in addition to starch (35 per cent), gum (± 7), pectin (17), iesin, wax, and

xx 80, 2400, xxu 456, 1156, 2322, 2829, xxu 404, 416, 2469, 2807, 2920, xxiv 2730, 3164, Arch Pharm, [8], xxv 329, xxvu 49, 211)

M Frennd (Ber, xxn 2829) suggests the following structural formules for hydrastinme and its derivatives:-

For hydrastme itself Fround suggests the following torunda -

$$CH_3 \begin{Bmatrix} O \\ O \end{Bmatrix} C_0H_2(CH_2 CH_3 NHM_0) C C C_6U_2(OM_0)_3 COOH$$

On decomposition into hydiastinine and opianic and, fission would take place at the point of triple linkage, both the acid and the basic derivative possessing aldebydic functions

¹ German; Kulumba or Columbo wurzel French, Racine de Columbo.

mineral matter (6 per cent.). Potassium nitrate has been found, but not tannin Berberine, the characteristic yellow alkaloid of calumba root has already been described (page 462)

Commung, or Cal um be Bitter, Capti, O, exists un calumber root to the extent of 0.34 to 0.40 que cant To atmest; the material is exhausted with boiling alcohol, the extenct evaporated to dryness, the residue taken up with hot water, and the filter of luquid staken with ether, or the tuncture is evaporated to a syrup, and shaken with chloroform. The chloroform solution is filtered, evaporated, and treated with 60 per cont alcohol, which dissolves most of the colouring matter. The residue is dissolved in strong alcohol, the solution decolorated with anuma charcool, and the columbin crystallised. Columbin is an intensely bitter, modorous, neutral body. It melts at 183, and crystallises from accite and solution in colourless trimetice prisms, very slightly soluble in cold water, more freely in hot

Columbin is sparingly soluble in cold alcohol, and in 40 parts of the boiling solvent. It dissolves with difficulty in cold ether, more readily in hot, and may be separated from betterine by sgitating the acidulated liquid with this solvent.

The solution of columbin is intensely bitter, it is not precipitated by tannin or any metallic salts

Columbin dissolves in strong sulphure and with orange colony, changing to deep red; on adding water brown fikers are disposited Columbin dissolves in aqueous sikalies, and is reprecipitated by scade On beating with causic alkin in and body is found According to Ho u d & columbin produces vomiting and distribund 10 19 gammes was futal to a fowly, death being preceded by diges-

tive disturbance and frequent evacuations (Pharm Jour, [3], xvi. 838)

Coursano Acup, C₂H₂O₂+H₂O, as prepared by treating the dired alcoholic extract of columba root with Imm-wates, and precipitating the solution with hydrochloric and it is a yellow sphorphous body, somewhat less bitter than columbin, nearly maschible in water, but thitle soluble in either, more resulty in acchic acid, and casily in alcohol

The alcoholic solution precipitates lead acctate yellow.

CAFFEINE AND ITS ALLIES.

Caffeine, the characteristic alkaloid of coffee, was obtained pure in 1821, when it was prepared almost simultaneously by Runge, Pelletier and Caventon, and Robiquet. In 1827, Oudry dusovered a similar principle in tea, and named it theire, Berzelius suggested the identity of this with caffeine, and this was afterwards established, as also was that of the alkaloid of guarane, called by Martius guaranine Madé, or Pranguny tea, and Koda naté contain the same alkaloid, while cocce contains the alkaloid theobromine (which may be regarded as a lower homologue of effective) in addition to small quantities of caffeine

Unlike the majority of the alkaloids hitherto described, theobromine and caffeine are not related to pyridine or quinoline They are respectively the di- and tri-methyl-derivatives of x anthine, C.H.N.O., a weak base forming the chief constituent of certain rarely-found urinary calculi, and existing constantly to a minute extent in normal urine and in most of the organs of the human body Xanthine itself is closely allied to uric acid, CaHaNaOa from which it differs by a single atom of oxygen, and from which it can be produced by treatment with sodium amalgam and water On adding silver nitrate to an ammoniacal solution of xanthine, an amorphous precipitate of the silver-derivative, C.H.Ag.N.O. is formed, and this when heated with methyl iodide is converted into dimethyl-xanthine or theobromine, C5H2(CH3)2N4O2. When the silver-derivative of theobiomine, C.HAg(CH.) N.O. is heated with methyl iodide to 160° C, for twenty hours, trimethylxanthine or caffeine, C,H(CH2),N,O, is produced.

The following formulæ show the constitution of caffeine and theobromins, and their relation to xanthine 1—

1 The formulæ given in the text are those proposed by Emil Fischer (Amaden, cexv 314) In the formulæ of Medicus (Amaden, claxv. 250), the CH and CO groups marked with an externsk are transposed.

Theophylline (see page 498), a base isomeric with theobromine, has been found in minute quantity in tea, as also has xanthine itself.¹

Caffeine. Theme Transthyl-xanthine Methyl-theobromine

CaHyaN,On, or CaH(CHa)aN,On.

The constitution and synthesis of caffeine have already been described (see page 473)

Caffeine exists naturally in the following sources, all of which are employed for food or preparing beverages —

a Coffee, 2 the dried seed of Coffee Arabica

b Tea, 2 the prepared and dued leaves of Camellia Thea.

o Maté or Paraguay tea, the dried leaves and twigs of New Paraguayensis.

d. Guarana or Brazilian chocolate, the dried pulp of the seed of Paulinia sorbits.

e Cola, the scods or nuts of the Kola tree (Cola or Sterculia acumunata) of West Central Africa

Casteine is found in other parts of these plants besides those commonly used for food, and also occurs in small quantity, together with the obromine, in cocoa

Caffeine can be isolated with facility in a state of considerable purity, but its quantitative determination is attended with considerable uncertainty, chiefly owing to the difficulty of completely extracting it from its natural sources (see page 448)

Caffeine is now prepared on a considerable scale from damaged tea ³ Several methods have been employed for the purpose, one of the simplest being to exhaust the tea with boiling water, boil

I For the mediation of matchine from tas, A. Baginaky effected the natural with datas subsime soil, instead the clean liquid with largita-with matchine with datas subsime soil, instead the clean liquid with largitary in excess, and then passed online drawled to procupitate the recess of faryta. After likewing and evaporating, ammoun and sulver mintar outside with subsidiary largitation from its solution in dilute nume and to which some once had been added. The northine silven institute obtained contained 3.5 dept. cent. of Ag. or very nearly the amount required by the formula C.HaN.O., AgNO. The weight obtained from 1 lb. of to was only 0.167 gramme (Farra Jose v, 38), ut at).

The absolute identity of the alkalouds of the and coffice is generally accepted, but cannot be said to have been established absolutely beyond doubt. According to faint of a Bit int in and Osa's (Pro. Bogd Society, 1887), this physiological effects of the alkalords extracted from ten and coffice withinform anised differences. These (from the appeared to be more powerful in itsortion than cafference from effects, and tended to produce hypthomical centractors of the voltancy nursices. These observations have not been confuned to the voltancy nursices. These observations have not been confuned.

³ A few years since the manniacture of cafferne was almost menepolised by Germany In consequence of a revised regulation of the English customs, the decocion with lithings or accitate of lead, and concentrate the filtered solution till the alkaloid crystallises out on cooling. The product can be punified by resublimation, or by crystallisation from hot water.

Cafferne forms long, whate, salky, floxable needles, which readily felt together to form light fleecy masses When deposited slowly from an aqueous or chloroformic solution, the crystals of cafforne present a characteristic appearance under a magnifying power of 100 to 300 danueters

It is generally stated that caffene crystallases from water with I arona (8:42) are out), but the proportion ordinarily present I norystallased caffene is sensibly less than corresponds to this formula respective and Lio big found 7.85 and Martius 8.14 per cent, and the author in two commercial spaciners obtained 7.05 and 7.10 per cent. It is probable that the deficiency is die officescence, for the water of crystallisation is lest by prolonged expositive over concentrated subjunize and at the ordinary temperature and pressure, so that the caffene so treated suffers in Turbio 1805 of weights 1100°.

On heating crystallised caffeine to 100° C the crystals become opaque and fraible, consequent on the loss of water, the residue consisting of analyshuse saffeine and dissolving without turbuildy in chloroform. According to Muilder, caffeine is deposited in anhydious crystals from alcohol or ether, and under cetain conditions from water also. It is possible that hydiation may depend on unrecognised conditions, such as those of temperature and concentration of the solution at the time of separation, and that commercial caffeine is a variable mixture of anhythous and hydrated crystals

Catterne does not evaporate with vapour of water, and undergoes no appreciable change of weight at 100° (A. H. Allen).² At 120° it volatilises very gradually, and at a higher temperature sublimes unchanged in long, silky needles.

according to which demaged ten as admitted duty-free, provided that its bent "densitued" and rendered wholly multi for hum in consumption by teach with line and assorteda, it has become possible to two such ten profitably for the manufacture of affirm. As a result, England has become the chalf set of the manufacture, and now experts the alkaloud to Genmany and America Appeared to the chalf of the chalf of

Mulder found 8 49 per cent of water, but that was by exposing the substance to a temperature above 120°, when more or less volatilisation must have taken place

² The statements respecting the effect of heat on caffeine are very discordant. According to A Wynter Blyth, eaffeine sublimes in minute needles at

- At 231°-233° C caffeine melts to a clear liquid, and at 384° (Strecker) boils and distris with partial decomposition, leaving no residuum
- 79° C , and volatilises completely at 120° . Other observers give much higher temperatures for its subliming point
- The behaviour of caffeine when heated has an important bearing on the methods of determining the alkaloid, and honce has recently been carefully re investigated in the author's laboratory by G E Scott Smith, C M Caines, and G S A Caines The following facts have been fully established —
- 1 Commsteal caffeine (crystallised) lost 6 9 per cent of its weight by prolonged drying over concentrated sulphune acid at the ordinary temperature and pressure.
- 2 Caffeins which has been dired at the cidinary temperature over sulphune and till constant in weight undargoes no further material loss on prolonged exposure in an open dish in the water-oven at 100°. The following results were obtained —

					Caff	etne	1	038
Weight	of commercial after long exp after heating	osure over H	804 at 9		1 000 g 0 931 0 929	, emmar	69p	er cent
,,,		10	61	19	0 929	15	71	22
,,	>>	10	51.	10	0 927	23	78	**

- 8. Notwithstanding the foregoing results, on hosting caffinise contained in a wish-fights, overall with nostlew statis-justs, over bulling water or not not of the water-oven for fifteen munitars, a distinct tilm appeared on the covering glass, and crystals of entires were tolsawable under the monoscope. The slight loss of weight observed when confines was exposed for many hours at 100° as doubbles due to relatifishing.
- 4 On exposing dry caffeine to a temperature of 120° in an air-bath, a very gradual but continual discusses of weight was observed, indicating sensible volabilisation of the alkaloid at the temperature employed. Thus —

					Weight of Alkaloid	L	DGB
					Grammos	Grammes	Per cont
foisture.	free	caffein	e talo	on,	0 9290		
fter hea	ting f	for 21	ours	at 120°,	0 9990	0 0030	0.82
	**	6		19	0 0270	0 0220	2 37
10		11	**	**	0 8008	0 0022	6 69
29	**	13			0 8314	0 0076	10 50
**	12	17	12	22	0.7850	0 1440	15 50
,	**	20	20		0.7054	0 1536	16 53
	**	24	19	10	0 7588	0 1722	18 68
2.7	**	20	**	12	0 7486	0 1904	19 62

Caffisine is odourless, but has a butter teate. It has a marked physiological action, and in accessive doses possesses decaded poisonous proporties. Administered to frogs, it produces tetanus and rigor of the voluntary museles. A cat was killed in thirty-five minutes by administering \(\frac{1}{4} \) gramme of lakalod. It all experiments with caffeine on the lower animals there has been increased frequency of the heart's extion, and repeated emptying of the bladder and intestines. After death, the alkaloid has been detected in the blood, the bile, and the urine. In man, offsition increases the heart's action, by stimulating the earlies museles, and excites the nervous system.

The British Pharmacopona gives from 1 to 5 grains as the medicinal dose of caffeine; the Genium Pharmacopona states the maximum single dose at 0.2 gramme, and the daily maximum dose at 0.6 gramme

The physiological action of infrisions of tea and coffee is in part due to the caffeine, but is largely modified by the other constituents, notably the tannin, extractive matter, and possibly the essential oil of tea, and the caffeel or essential oil of coffee.

Caffeine is only sparingly soluble in cold water (75 to 80 parts), but tolerably readily in hot (10 parts). It dissolves in about 35 parts of cold rectified spirit, but it is much less soluble (1 155) in absolute alcohol. In cold ether it is very sparingly soluble, more readily in anythe alcohol, chloroform and bonzene, but nearly

- 5. Onffenns which had been recently sublimed and was consequently anhydrous, melted at 251.5°C, and resolution at 225°C. Streckor gives the making-point of anhydrous cuffence as 224°, and Bredermann at 250.0°. Mulder gives the melting-point at 177.3°, which is outsimly too low.
- 6 Caffeine which had been recordly sublimed and then dissolved in water, acticolo, ethin or chiroform, in sech case left the original weights of alkaloid on evaporating the solution and exposing the issuine at 100° The same result was obtained with recently-fused outliens. As sublimed and fused caffeine are cuttainly subjectors, it follows that the alkaloid left on evaporating its solutions in the above volvents is also analytics.

7. When a known weight of caffines was repeatedly instead with a small quantity of wast, and the luquid expented to drayses at 100°, the original weight was always recovered. When caffines, parviously dued at 100° or 120°, or recently sublumed or fenced, was densired in 1000 are of catalities water, the solution concentrated by boiling over a naked flame, and the exporation completed in a platinum dish at 100°, the reaches being finally fixed in the water-own, the weight of sikeland originally taken was afriedly recovered. This proves that coffens does not volatilise with steam during the exporation of its solutions (A. H. All in J. Zham J. 2007, 133; Xun. 213).

msoluble in carbon disulphide and petioleum spirit. Chloroform and benzene dissolve out the alkaloid even from its acidulated aqueous solutions, but the agricultons must be several times repeated to effect complete extraction

Concentrated sulphure and converts cafferne into the sulphate, but does not colour or otherwise change it even at 100° C 2

Hydrochlous and has no actom on cattenne bolow 200°, when heated unden pressure with concentrated hydrochlora can be 280° for any to twelve hours caffenne yields ammouns, methylamine, sercesine, carbon dioxide, and traces of formes and . The volume of methylamine produced is double that of the ammouns, which proves the presence of three NMe groups in caffeane, and establishes the following formath for the reaction $-C_0 \frac{11}{10} N_0 Q_+ + 6 H_1 O_1 = N H_2 + 2 N (CH_0) H_1 + C_1 H_1 N O_2 + C H_2 O_1 + C O_2 (E S ch m it d, Amadea, exciv. 1270) s^2$

When caffeine is warmed with dilute caustic alkali or boiled with concentrated baryta-water, it at first assimilates the elements of

1 A Commaille (Compt. Rend., exxx. 817., Jose Chem. Sec. xxx. 778) gives the following figures for the solubility of hydrated and anhydrous caffains in different menstrus —

	Parts of Solv	ent required fo	z 1 of Cufeine
Solvent	At 15°	to 17°C	At Boiling point of Solvent *
	Hydrated	Anhydrous	Anhydrous
Wates, Rectified aparts, Absolute alcohol, Commercial esher, Fure analysions other, Chiloroform, Carpon disulphild, Fetroleum esher,	08 40 476	74 41 165 526 2238 77 1700 4000	2 2 32 277 51 220

" The hot water was at 85" only, not at the boiling point

⁸ Experiments by the author aboved that pure caffune was wholly unchanged when hoteld in the wates over fine soveral hours with concentrate all subhume and On dissolving the product in wate, boiling with earlie of lead, filtering, and cat testing and extit esting with children's the original weight of caffine was recovered. Some samples of commercial caffune darken alightly when heated with subjusture and

Schradt thought it possible that theologimus might be formed in this reaction by the climination of a methyl-group, but was not able to detect it. The methylamino was separated and punified by convenion into the chloroplatinate. The saucosine was identified by means of its coppose salt.

water and is converted into an acid containing $C_8 II_{12} N_i O_3^{-1}$ On further treatment, this body splits up with great facility into earbon dixide and the base caffe of $1n \in O_{\rm FR} N_i O_3^{-2}$ On still further boiling with the alkali this is again decomposed with formation of carbon dioxide, formic acid, ammonia, methylamnie, and sercome (methylamnie, and sercome (methylamnie, and sercome (methylamnie)).

The author has proved that eafleme readily undergoes decomposition when boiled with lime-water, a fact which has a practical bearing on several of the published processes for its determination. When eafleme is boiled with magnessa and water, the decomposition is instantional, and with litharce there is no change.

CAFFEIDINE CARBOXYLIC ACID, CaH12N4O3, or CyH11N4O COOH, is best prepared by digesting finely-divided cafform for some hours at 30° C in a dilute solution of caustic notical or soda, nentralising with acctic acid, adding curric acctate (avoiding excess), and decomposing the resultant precipitate by sulphuretted hydrogen. The liberated acid obtained on evaporation of the filtrate to vacuo at the ordinary temperature, may be purified by solution in chloroform and meanitation with benzene, and is thus obtained in the form of a thick oil, which on exposure to the au solidifies to a yellowish-white. slightly envetaling mass, your soluble in water to a strongly and bound. It is soluble in alcohol and chlorotom, but insoluble in benzene. On boiling the aqueous solution of enfferdme-entboxylic acid, carbon dioxide is evolved and a reddish oil remains, which when started up with a small quantity of sulphure acid and treated with alcohol solidifies to a white crystalline mass of caffeeding sulphate The reaction affords a ready method of preparing caffednie. It is merely necessary to decompose the copper salt with sulphinetted hydrogen, evaporate the filtrate rapidly, and treat it with strong sulphuic and The copper salt of caffending-carboxylio acad, Cu(CaH11N2Oe), 19 a pale blue civitalline pewder, nearly insoluble in water and wholly so in alcohol. The burnum, calcium, zino, cadmium, and magnesium salts are nearly insoluble in water, but the lend salt is soluble KA is a yellow oil On adding mercune chloride to the solution of a soluble caffedure-carboxylate, a copious white precipitate is obtained which appears to contain (CgH11N4O3)allg,2llgCla It this be suspended in water and decomposed with subburetted hydrogen. the filtered hand leaves enflexing hydrochloride on evaporation

**OAFEDITA'S, GIJLANO, may be obtained as above described, or may be operated by louing caffines with a solution of 10 pairs of rystallane-ling repeated by louing caffines with a solution of 10 pairs of rystallane-ling free half on hour, or until ammons and multylenume begin to be evolved by excluding the fitteed thank with dutte onlybune scal, and evaporating the fitteed hand with dutte onlybune scal, and evaporating the fitteet of them strying a likeline lund, reality-solution in water, school and chlorofum, but with defineding me disk if reduces value or aske, even in the cold, and decomposes very readily into ammons, methylamon, and choles it re plan of (amethylamon and C, Jajla-Ko, No.). Caffeditum intrate, hydroloumide, and hylrochloride or spitallas well B (1470/G, systallas less from water in large compacy vider woulds.)

When cassen is heated with sode-line to 180° , ammonis is volved, and on borate and a large quantity of c y an 1d of formed According to R of R of R of R of this last product distinguishes cassen from a pignated with excess of sode-lines, the introgen is ordived as animons, any cyanide formed as an intermediate product at a lower temperature being decomposed in the usual manner, but in rules to ensure complete conversion of the nitrogen into animons, it is better to mix the cassen of cassens of the conversion of the nitrogen into animons, it is better to mix the cassen of the conversion of the nitrogen into animons, it is better to mix the cassen of the nitrogen into animons, it is better to mix the cassen of the nitrogen into animons, it is better to mix the cassen of the nitrogen into animons, it is better to mix the cassen of the nitrogen into animons, it is better to mix the cassen of the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons, it is better to mix the nitrogen into animons and the nitrogen into animons.

When caffine is treated with bromine-water, avoiding excess and the liquid evaporated to dryness at 100°, y selfowsh receives left, which becomes crimson-ied on further hesting, and is turned a migraficent purple by ammonia. The reaction is very identicate, and is not affected by a considerable excess of ammonia On adding causies code complete and instant decolorastion occurs.

Another modification of the test consists in treating a munute quantity of the bold substance (such as a residue) of caffune left on evaporation) in a porcelain dush with a few drops of strong hydrochloric acid and a minute crystal of potassium chloric, and ovaporating the highit to dryness at 100°. When cold, the reddish-y-allow or pinkish resultes is cuntiously meastened with ammona, avoiding an excess, when the characteristic puiple coloration is produced, or, preferably, it is exposed to ammoniacal vapours by inverting the dish bearing the resultee over another containing stong ammonia

The products of the exidation of cafeine melinde am alice act d_1^{-1} which by subsequent treatment with ammonia is converted into murexolm, the sections being identical to the eye and parallel in chemical change to those yielded by uncaerd under like conditions. Thus—

Strong nitric and may be substituted for the bromine-water or hydrochloric and and potessium chlorate, but the reaction is in that case far less distinct and easy to regulate, and excess of ammona must be emefully avoided.²

¹ AMALIO ACID forms colourless crystals which stain the skin red, and are yearingly soluble in water or alcohol. It reduces silver salts, and forms deep violet compounds with potash, sods, and bayta

⁹ O. Hehnos, in a private communication to the author, points out that, if the altric and used be perfectly pure, caffeine fails to give the mirrexon reaction, but that in presence of a minute trace of hydrochloric acid the colour is readily developed.

Theobromine and xanthine give similar reactions to caffeine with an oxidising agent and ammonia. The purple colorations due to caffeine and theobromine are decolorised by adding caustic alkali solution, but that due to uric acid is changed to blue.

When cafferne is heated with a large excess of nitric acid, it is converted into cholestrophanel or dimethylparabanic acid, Ca(CHa), NaOa a body which crystallises in pearly laming. melting at 1455°, boiling at 275°-277°, and difficultly soluble in cold water and alcohol It is decomposed with great facility by alkalies into symmetrical dimethylures (incling at 97°-100°) and oxalic acid. Hence on adding ammonia and calcium chloride to its aqueous solution, and warming the liquid, calcium oxalate is precipitated.

Cholestrophane is also produced (354 to 418 per cent) by oxidising caffeine with chromic acid mixture, the main reaction being -

 $C_0H(CH_0)_2N_4O_2 + O_3 + 2H_2O = C_3(CH_3)_2N_2O_4 + NH_2(CH_3) + NH_2 + 2CO_3$

Caffeine is very imperfectly precipitated by the usual alkaloidal reagents. No reactions result with iodised potassium iodide and Mayer's solution, which behaviour distinguishes caffeine from nearly all other alkaloids except theobromme and colchience Potassiobismuth iodide precipitates caffeine after a time from moderately dilute solutions (1 8000) Phosphomolybdic acid produces a yellowish precipitate, soluble in warm sodium acetate solution, the liquid depositing free caffeine on cooling (CaH10NAOa HCl)aPtCl is obtained on adding hydrochloric acid and platinic chloride to a highly concentrated solution of caffeine, as an orange precipitate soluble in 20 parts of cold and an even smaller quantity of warm water, crystallising again on cooling

A solution of caffeinc in 200 parts of water gives an immediate and abundant precipitate on adding a saturated solution of mercuric chloude With a more dilute solution (1 1000) crystals appear in a few minutes, and in an hour or two an abundant crop of large acteular crystals is obtained. With a solution of caffeine in 4000 of water crystals appear after a few days. The precipitate contains C₈H₁₀N₄O₂,HgCl₂, and is much less soluble in excess of the reagent than in pure water Hence the best results are obtained by adding an equal measure of a concentrated solution of mercuric chloride to the liquid to be tested. The compound is soluble in about 260 parts of cold water, and more readily in hot, crystallising out again on cooling. It also crystallises from hot alcohol The

¹ The name cholesticphane is due to Stenhouse, and has reference to the resemblance the crystals have to cholesterin (Vol II page 312). VOL III PART II

compound is not sufficiently insoluble to be applicable to the quantitative precipitation of cases (R. H. Davios, Pharm Join, [3], xxi 253)

Gallotannic acid precipitates moderately diluto solutions of the caeffance, the precipitate being somewhat solutible in excess of the reagent A difference of a few degrees in temperature greatly alters the solutibity, and hence a solution of properly alphase strength may be perfectly limput at one temperature, and become completely opaque from separation of amoujhous caffeing sellocations of the cause of an influsion of tea becoming turbid on cooling

SALTS OF CAFFEINE

Caffeine is a very feeble base. Its aqueous and alcoholic solutions have no action on litmus, and it is extracted from aqueous liquids by benzene and chloroform, even in presence of a fiee mmeral acid This behaviour is doubtless due to the facility with which the majority of caffeine salts are decomposed on dilution They are decomposed by alcohol and other as by water, and the salts with volatile acids (eg, acctac) are decomposed on exposure to air The hydrochloride leaves meiely free caffeine on exposure to 100° C The author found that on adding free eafferne to hot water containing a trace of sulphuric acid and coloured with methyl orange, the red colour of the liquid was immediately destroyed. proving neutralisation of the acid, but an acid leaction was 10established when standard acid had been added equivalent to only about alout of the eaffeine present Owing to these facts, certain devises have to be employed for the preparation of the majority of the salts of caffeine The oxalate 1 and salicylate are sparingly soluble, and can be readily prepared by mixing equivalent quantities of the acid and alkaloid in aqueous solution. The estrate is best obtained by mixing a chloroformic solution of caffeine with an alcoholic solution of citric acid, and evaporating the mixture to a syrup When molecular proportions of eaffeino and a mineral acid are mixed together in presence of oxeess of water, no combination ensues. If the quantity of water is insufficient to dissolve the alkalow, the latter remains suspended in the liquid in an unchanged condition If the hquid is allowed to evaporate spontaneously, the acid ultimately becomes sufficiently concentrated to act on a portion of the caffeine, and a true salt crystallisos out, intermingled

Caffeine exalate is said by Leipen to be an exceptionally stable salt. It can be increptablised from water, but the author found that the whole of the caffeine could be removed by chloriform from an aqueous solution containing a considerable excess of exalic and.

with cystals of the unaltesed alkaloud. But as the acid is weakened by its combination, the formation of the sait is setaided till further concentration has taken place. Hence the change is progressive and continuous, the coffeine guadually dissolving and again cystal-lises out as a sait, though at the very last cystals of the uncombined base can be observed in admixture with the increasing crop of time sait. Dy employing a considerable excess of north the processing greatly lastened, and a product free from uncombined alkaloud sobtamable. With an excess of said, and a sufficient degree of concentration, the alkaloud will momentarily dissolve to a clear solution, and then almost immediately crystallise out as said.

The foregoing observations are due to H W Snow (Phanm. Jour, [8], xxi, 1185), who gives the following as the composition

of the principal salts of caffeine -

Caffeine hydrochlomde, B,HCl+2H₂O Caffeine hydrobromde, B,HBr+2H₂O Caffeine hydrobromde, 5fB,HNO $_2$ J+4H₂O Caffeine sulphate (normal), B,H $_2$ SO, Caffeine sulphate (normal), B,H $_2$ SO, Caffeine sulphate, B,H $_2$ C,O $_1$ Caffeine salleplate, B,H $_2$ C,O $_1$

Caljeve hydrochlo ide crystalluses in colouless prisuatic needles it loses the whole of its cut at 75° C. The sulphate is adopted from a hot alcohole solution in simming needles unchanged at 100° Cafficientist afforms fine transparent cystals, which when dropped into water become opaque, and are converted into pseudomorphs consisting of mucoscoure needles of free caffine.

Cadjane citrate is official in the British Pharmacopress of 1885, where the formula CaH₂M₂O₂M₂CaH₂O₃ is assubed to it. The BP. exticle is generally regarded as an indefinite, unistable, inaccurately described, and superfluous preparation (Pharm Jour, [5], xiz 252). Free caffeine has not unfrequently been sold as the citrate. The proportion of acul can be directly assentiated in the citrate and other caffeine sails by straining the solution with a standard castical callul (or prefaibly baryls and phenolphithalem, and the total caffeine can be molated by agitating the neutralised or original aqueous solution with chloroform. On treating the div substance with cold chloroform, only the uncombined caffeine, if any, will be dissolved out (J U I Lo y U).

A strong and stable solution of caffeine can be readily prepared by dissolving it in benzoate, cinamate, or salicylate of sodium or ammonium. Such solutions are employed for hypodermic nigetions, and caffeine phenate and phthalate have been applied to the same purpose. ISOLATION AND DETERMINATION OF CAPPRINE

None of the compounds of cafferne are sufficiently stable or insoluble to be of service for the separation or precipitation of the alkaloid, which is always determined by weighing it in the free state The isolation of eaffeine presents no difficulty, and may be effected by a variety of methods. The majority of these depend on the treatment of the substance or its aqueous infusion with lime, magnesia, lithauge, or basic lead acetate, to render the tannin, &c, insoluble, and crystallisation of the caffeine from the concentrated filtrate, or extraction of it by benzene, other, or chloroform To ensure the absence of morganic salts, the alkaloid should be sublimed or shaken out from its aqueous solution by chloroform Provided that the caffeine isolated be well crystallised, colourless, free from acid or alkaline reaction to htmus, completely soluble in chloroform, exerts no reducing action on Fehling's solution, and leaves no ash on ignition, it may be regarded as pure

Although the mointon of caffeine in a state of absolute purity may be easily effected, the accurate determination of the proportion of alkaloid present, especially in tea, is attended with great difficulty, and hence most of the published results represent the proportion of caffeine isolated, rather than the amount existing in the substance examined. When once in solution, several methods may be used, though even in this case some of the published processes give results which are vory gravely wide of the truth. As consequence, the great majority of the published determinations of affence are completely workless, and even where a number of figures have been obtained by the same process. Whey do not necessarily became any definite relation to each other

The determination of the alkaloid in tea has recoultly been the subject of a very large number of experiments in the anthor's laboratory by C M Causes, G S A Canses, and G E Scott Smith (*Phas m Jour*, [3], xxiii 215). The following facts have been fully established —

I Aqueous solutions of caffeine, even when very dilute, may be concentrated by boiling, and subsequently evaporated to dryness at 100° without the least loss of alkaloid (see page 477)

2 Caffeine may be completely dehydrated at 100° in the wateroven It undergoes no appreciable loss by volatilisation when exposed to 100° for many hours, but sublimation to a minute extent can be proved by the aid of the microscope (see page 476)

Cafferne cannot be estimated, even approximately, by crystallisation from water, the amount which remains obstanately in solution, in the presence of saline matters, often exceeding that which can be separated as crystals

- 4 Caffoune can be completely extracted from its accludated or slightly ammonated apresses solutions by repeated agitation with chloroform. In the author's experiments, from a solution algibilly accidently as a solution algorithm of the solution algibility accident with sulplumo and, the first treatment with chloroform or similar and the solution of the alkaloid, but it is desamble to agitate a fifth time and evaporate the separated solvent apart, to prove that no more caffene is being dissolved. In this last case, the solution may be advantageously rendered ammoniscal, or a loss of 0.001 to 0.00 gramme of caffene way occur, puolably owing to the existence of traces of caffene sulphate, sepseally where the solution is strongly calculated with sulphuro acid. On distilling the chloroforms solution of caffeine, and drying the residue at 100° C, the alkaloid is obtained in a perfectly analytous condition.
- 5 Chancol cannot be employed for decolorising caffeine solutions, without a considerable absorption of alkalod, which is retained with extense persistency. If the caffeine isolated be colored, it may be dissolved in a little hot water, and the fitted solution evaporated to dryness, but there is little difficulty in isolating the alkaloid in a sonw-white condition.
- 6 Caffeine is completely unchanged by leating to 100° with strong hydrochloric acid, or with sulphune acid diluted with onething of its measure of water. On treating the mixture with water, the whole of the alkaloid may be recovered by agitation with chloroform, as in 4.
- 7 Caffeine is readily decomposed by alkalies By warning with dilute causic soda, it easily undergoes change, and by boiling with lime it is partly decomposed, with formation of ammonia and methylamine (see page 479)
- 8 When commercial caffence is treated with sguited magnesia and water, and the mixture distilled, a slight but distanct formation of ammonia is observed, apparently accompanied with timees of volatile ammers. But the volatile leases are found chiefly in the first fractions of the distillate, the latter portons being quite free employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other volatile bases is employed, the formation of ammonia and other other bases of the latter case the production would continue throughout the distillation. On filtering from the magnesia and extracting the filtrate with chloroform, the original weight of caffeine can be recovered, if the pure alkalord was originally sunployed.

- 9 If a mixture of caffeine with magnesia be made into a paste with water and dried, the alkaloid can be wholly extracted from the mixture by prolonged treatment with chloroform
- 10 When one part of caffeine is dissolved in hot water, and a solution of two parts of galloidamin each added, the caffeine can be securitely determined by precipitating the solution with lead acctate and extracting the concentrated filteriae with chilosofum If the higher becomes the first the concentration of the work of the magness, and dried at 100°, the whole of the alkknloid cannot be extracted by boiling the powdered mass with dry chiloroform, however long the treatment be continued If taming repeared from tea be substituted for galloidamic soul in the foregoing experiment, a similar result is obtained.
- 11 When a decoction of tea is substituted for the foregoing artificial mixture of caffeine with excess of famini a precisely similar result is obtained. Whether said or magness be used, the alkaloid is only partially extracted, even after prolonged boling with chloroform or either ⁷ Thus, decoctions prepared by
- ¹ The following experiments were made by G. E. Scott Smith in the author's laboratory. Fifty grammes weight of commercial black tea of medium quality was produced and boiled with waton for threty immits. The solution was filtered and made up to I litre after cooling. Abunot parts of the solution were then tested in the following manner.
- A 100 c o (-5 grammes of tea) was evaposated to a symp and mixed with 5 grammes of ignited magnessa. The mixture was dired tho oughly at 100°, powdered, and boiled with ether froe from alcohol and water

```
Cofficine orticated by 6 hours' treatment, 0 0009 gramme.

10 000 gramme.

10
```

On subsequently boshing the residue with alcohol an additional 0 0605 gramme of caffeine was extracted, making 2 59 per cent in all

- B. Was conducted like A, but dry chloroform was substituted for other The total cafferno extractable by chloroform was I 54 per cent
- C Conducted like A, but rectified epart was employed at once It extracted 28 Iper cent of became a callone, which was reduced to 2.78 per cont by re solution in water and extraction with chiptorform.
- D. Conducted hire R, but sand was substituted for magnessa Treatment with dry chalonform extracted successively 0.0846, 0.0175, 0.0135, and 0.010 gausses of calicus during mue hours' treatment. On subsequent treatment with sloched much tannus and colouring matter was extracted. This was promptated by lead acotate, and the concentrated fiftstate shaken with chloro-form. Additionally pried, 0.070 gamman, sunking a total yeal of 2.072 per California. Additionally pried, 0.070 gamman, suchage a total yeal of 2.072 per California of the cafferes but not the whole should be extracted by chloroffer in the absence of magnesses not or ordered.
 - E 100 oc (-5 grammes tea) was heated to boshing, treated with solid

looling two separate samples of black tea with water were each druided into two equal parts. One of these was preceptated by lead acetate, and the enfirme recovered from the filtered and concentrated liquid by repeated agriation with oblicoform. The other halves were componented to dryness with magness and the powderreadure thoroughly exhausted by boiling with chloroform, and subsecuretity build with alcohol for a lone time.

		Sample A 30 Minutes' Bolling	Sample B 20 Minutes' Boiling
Lead process,	,	8 31 per cent	2 07 per cent
Magnesia process by chloroform, , ,, by alcohol, .	•	118 ,,	0 90 1 16 2 98 per cent

In other experiments with mixtures of caffeine, tea-tainin, and excess of magnesia, from 8 to 10 per cent of the alkaloid was not extiactable either by chloroform or alcohol, but could be recovered by teatment with water

- 12 When finely-powdered tea as mixed with elaked him, upitied magness, or sand, made into a paste with bot water, and the mixture thoroughly dired at 100°, only a fraction of the total alkaloid can be extrahed with elhoroform, however carefully the process be conducted. On subsequently teating the mixture with alcoloi, the greater part of the remaining eafforms is ultimately insolved, but prolonged treatment by boiling alcohol is necessary to extract the caffeine from a mixture of tea-extract or powdered tea with magness, and complete extraction is always doubtiful
- 13 When a decochon of tea is treated with base or neutral scotate of lead a voluminous prepriate is formed. If an alquot part of the liquid be filtered, concentrated, and treated with suphurested hydrogen, sulphurous acid, sulphure acid, or sodium phesphate, to remove the excess of lead, and again filtered, the caffeine may be extracted in a condition of perfect whiteness and purity by agitation with chloroform.

lead acetate, filtered, and an aliquot part of the filtrate concentrated, freed from lead, and shaken repeatedly with chloroform Caffeine was recovered equivalent to 2 68 per cent of the tea

¹ The remarkable fact of the telenton of the caffeine of tax by line or magnesian in form incompletely disorbed by cholorious was first observed by B H Paul and G E Scott Smith (Rhom. Jose, [3], xx 832). Little more than one thuil of the total caffism was orticatible by chlorious information with the winking and the more than one-half from the magnesia mixture. Psy subsequent treatment with abool the renaming caffisme was disorbed.

14. By prolonged boiling with litharge a decoction of tea becomes completely decolorsed, but the process is tedious If after a time a small addition of lead acetate be made, clarification occurs in a few immutes, and an aliquot part of the liquid may be filted and instant of an interest of an interest of the second of the contract of a process.

From the foregoing statements (10, 11, 12, 13) it, is wednet that the determination of coffinms when in a state of solution presents no great difficulty, though the windly-word plan of evaporating the lought with sead and lime or magnetis, and extracting the disciding statement of the statement of the disciding statement of the registration of the alkadout is retained by the registration these, as fact which suggests that it exists pathly in some modulute combination only gradually decomposed by boiling water or alcohol.

On the supposition that the cellular structure of the tea is the cause of the obstinate retention of the caliform, Z o ll or (Zwisch Anal Chem, xii 106) has proposed to treat the finely-powdered tea with strong sulphurae and duluted with one-third of its

¹ The following figures, obtained in the author's laboratory, show the rate of exhaustion on treating powdered black tea with hot and cold water —

Cafferne extracted t	y Cold Water	Cafferns extracted by Borling Water				
In 3 days, Additional 2 days,	1 81 per cent 0 29	In # hour, In additional 2 hours.	2 46 per cent			
" 2 days,	0.70* ,,	,, 4 hours,	016 ,,			
,, 6 days,	013 ,,	,, Gridars,	0 01 ,,			
Total in 19 days,	8 15 ,,	Total in 12½ hours,	3 85			

* These two figures have not been transposed

Thus the extraction of the caffeine by boiling water was practically complete after 6 hours' treatment, while with cold water the total amount was not dissolved after 19 days' treatment

In both the hot and cold water experiments, the infusion reduced Fehling's solution after removal of the tannin by lead accepte. The caseme did not reduce the copper solution either before or after boiling with drive acid

² The author has proved the presence of a glucoude in some teas

measure of water, and heat the maxture at 100°, tall the cells are thoroughly broken up. Some water is then added, an excess of hydrated oxide of lead stirred in, and the mixture dired and exhausted with alcohol of 86 per cont. The alcoholic solution is disclorized with animal charcoal, and evaporated till caffeine crystalliaes on cooling. From the mother-luque, the residual caffiens is extacted by ether. Coller obtained the high proportion of 492 per cent of caffiene from a high quality of Himalayan tea, in addition to an anytexable quantity of theorbromine

The author has made a number of experiments on the lines of Caller's process, modified in various manners, but, chiefly through the remerkable persistency with which caffeins is absorbed and retained by the earbon formed by the such teastment, they are not hitherto resulted in the evolution of a practical analytical method?

1 On treating powdered tea with slightly diluted sulphure acid, and heating the mixture in the water-oven for an hour or two, a black product is obtained which powders readily On boiling this product with water, a perfectly colourless solution is obtained, from which, after concentration, perfectly colourless cafferne is extracted by agritation with chloroform, cither with or without previous iemoval of the sulphings and by boiling with lithange or white lead, or neutralisation with ammonia. The fact that a colourless liquid is obtained on treating the charred tea with water is due to the absorption of the colouring matters by the finely-divided carbon formed. Unfortunately, this product also takes up a considerable proportion of the caffeins, and retains it with such obstinacy that it is only extracted by prolonged and reneated treatments with alcohol Although the entire amount presont is ultimately obtainable in solution, the extraction is too uncertain and tedious to render the method a desirable one in mactice. Exhaustion direct with alcohol, ethor. chloroform, benzene, or water, either with or without provious neutralisation of the and with litharge or magnesia, equally failed to cosine ready extraction Of the numerous experiments made in this direction the following may be mentioned. Twenty five grammes of ordinary black tea of medium quality was finely powdered, and treated with 10 c c of sulphmic and diluted with one fifth of water The mixture was heated at 100°, treated with a hitle water. and ground with excess of litherge till noutral. The mixture was reduced, and thorongily exhausted successively in a Soxhlet-tube with boiling rectafied spirit, boiling proof spirit, and boiling water. The solutions were evaporated, and the cafferne extracted by repeated agritation with chloroform. The following were the results obtained -

		Yield of Caffeine
By strong alcohol (sp. gr. 838),		3 03 per cent
By subsequent treatment with proof spirit,		0.50 ,,
By subsequent treatment with water,		0.21 ,,
Total.		374

The caffeine isolated was snow-white. These results show that the alkaloid is unaltered by the treatment, and if extraotion could be effected with cortainty

As the result of very numerous experiments, the author gives the preference to the following method of extracting and determining the caffeine in tea. It closely resembles a process employed by S t a h lschmidt (Chem Centralblut, 1861, 396) -Six grammes of finely-powdered tea is treated in a flask with 500 ce of water, which is then kept boiling under a reflux condenser. No Soxblet extractor or similar arrangement is so effective or iapid as actual boiling with the water Alcohol effects no quicker or better extraction than water, and has the disadvantage of dissolving chlorophyll. After six or eight hours' boiling, the decoction may be filtered, the residue washed on the filter, and the filtrate made up with water to 600 e.e. It is then heated nearly to boiling, and about 4 grammes of acetate of lead in powder added, a reflux condenser attached, and the haud boiled for ten minutes. If on removing the source of heat the precipitate does not curdle and settle readily, leaving the liquid colourless, or nearly so, a further addition of lead aectate must be made and the boiling repeated. When elarification is effected. the liquid is passed through a dry filter. Five hundred c.e. of the filtrate (= 5 grammes of tea) is then evaporated to about 50 ec. when a little sodrum phosphate is added to precipitate the remaining lead The liquid is filtered, the precipitate washed, and the filtrate further concentrated to about 40 ce, when the caffeine is extracted by repeated agricultanes with chloroform, at least four treatments with which are necessary to ensure the complete extraction of the alkaloid 1 The separated chloroform solutions are mixed, and distilled in a tared flask immersed in boiling water. The last traces of chloroform are removed while the flask is still hot by a current of air, and the residual alkaloid is weighed. The caffeine thus isolated is snow-white in colour, neutral in teaction to litinus, and completely volatile and soluble in water It does not reduce Fehlung's solution either before or after boiling with dilute acid

As a precention, the exhausted tea-powder should be again bouled with water, and the decotion treated as before When experience has proved this to be imnecessary, the piecess can be shortened by boling the tea with 600 c of water in the first place, and adding lead acctate without previously filtering from the schausted tea. This modification becomes necessary in the case of schausted tea.

by a engle solvent, the process would possess marked advantages. Substitution of magnesia for the oxide of lead, and various other modifications of the details equally failed to give a satisfactory result.

In the great majority of cases the chloroform separates readily. Should an obstinate englision be formed, the best plan is to place the mixture in a fask, distil off the chloroform, treat the residual liquid with a few drope of basic acetate of lead, filter, and agitate the filtrate again with chloroform.

certain teas (e g , gunpowder), the aqueous decoctions of which filter very slowly

The following neutles by the above process were obtained by G M G an ne in the author's laboratory (Phow M dow, [3], xxii. 218) In some instances the eaftene extracted by half as hour's bolinique was determined, in addition to the total amount obtained by ax hours' loohing with water. The results refer to the mostunctree teas, which were represe native commercial samples —

		Caffeine,				
Description of Tea.	Tanuin , by Load Acctate	Extracted in 80 minutes	Total, extracted in 6 hours			
Ceylon, whole leaf (Pekce),	13 01 per cent	\$ 40 per cent	3 85 per cent.			
Coylon, broken leaf, .	1231 "	ì	4 68 ,,			
Assam, whole leaf (Peloe),	10 08 ,,		4 02 ,,			
Assam, broken leaf,	11.33 ,,		4 02 ,,			
Java Pukoe,	12.98 ,,		875 ,,			
Kaisow, red leaf, .	11 35 ,,]	841 ,,			
Moning, black leaf,	1176 ,,	844 ,,	874 "			
Moyune Gunpowden,	12 95 ,,	276 ,,	2 89 ,,			
Natal Pokoe Souchong,	900 ,,	271 "	8 08 ,,			

The foregoing piocess is applicable to the determination of the caffinion in coffee, of which 12 grammes may be conveniently employed. In the presence of choory the extracted alkinoid is hable to be strongly coloused, in which case it should be redused with water, a few drops of caustic soda added, and the liquid again exhausted with chloroform

An alternative process for the determination of casilene in feas is that of P a u I and C ov n is $P(Ram\ Mou\ P)_3$, xmu 147), which in some respects resembles a method described by V or n in n in $(Aich\ Pham\ n, [2], 1xviii 148), and with centaun modifications communicated to the author by <math>A$ J C ov n if v is as follows —Five grammes weight of finely-powdered tea is well mixed in a mortar with 2 grammes of ignited magnessa, the mixine thoroughly mostened with hot water, again triturated, and then those at 100° it is next extracted with bolling alcohol, l and then clined at 100° the size v is in v in v

³ Experiments made in the author's laboratory showed that even with the most careful treatment it is difficult to casure o m_i-lete extraction of the caffeine, a small additional quantity being subsequently obtained by treat ment with water. until sxhausted ¹ The united chloroform solution is then agitated with a very dilute solution of caustic sods, which removes a little colorung matter, so that on subsequently distilling off the chloroform in a weighed flask, the caffeine is obtained perfectly pure and coloriless, or at most with a faint green large.

By the foregoing process, Paul and Cownley (Pharm Jour, [3], xviii 417) found Indian and Cingalese teas to contain a much larger percontage of caffeine than, owing to the faulty methods of analysis employed, is commonly supposed. The proportion of alkaloid isolated from commercial samples of all qualities, and containing from 3 6 to 6.8 per cent of moisture, ranged from 3 22 to 4 66 per cent on the team its commercial condition (equal to 3 57 to 4 99 per cent in the moisture-free tea), and boic no relation to the so-called "strength" of the tea Jave tea approached Ceylon tea in the proportion of caffeine present (294 to 378 per cent), but China and Japan teas were generally poorer in alkaloid, the proportion in these products minging (for a limited number of samples) from 2 20 to 3 46 per cent J. H S mall obtained, by Paul and Cownley's method of assay, from 179 to 230 per cent of cafferne from Japanese teas, and from 2 38 to 3 54 per cent from Chinese and Indian teas.

P au 1 and C o w n l a y have also employed the foregoing method of determining caffeeine for the assay of coffee (Pharm Jon. [3] xvi 565, 643) The caffeine obtained by ovaporation of the oblivoriom is hable to contain a small quantity of a hownish waxy or resnous impurity, and hence should be purified by resolution in boiling water, and recovered by evaporating the inflatered solution and drying the residual alkaloid at 100°. By this process they found the proportion of caffeine in coffee-berries to vary within companitively narrow limits, and not to be materially affected by reasting Hence they recommend the determination of the alkaloid in commercial coffee as a neems of estimating the proportion of choory or other admixture present

Theobromine. Dimethyl-xanthine.

The constitution and synthesis of theobromme have already been described (page 473). It is the lower homologue of caffains, to which alkalord it presents a close general resemblance, but differs considerably from it in its solubilities.

³ In Paul and Cownley's experience, six or seven snocessive treatments with chloroform (using from 30 to 40 c.c each time) are necessity to effect the complete extraction of the caffeine from the solution yielded by 5 grammes of tea.

Theobiomine is isomeric with theophylline and paraxanthine Theobromine exists naturally in eocoa, the seed or bean of Theobioma cacao, and together with caffeine in the kola nut (Steroula acummata) An alkaloid apparently identical with theoformine was found by Zoller in a specimen of Himslayan tea.

Theobromine forms a white, crystalline powder, which under the microscope appears as trimetric needles and club-shaped groups When heated to about 290° it sublimes without decomposition or previous fusion.

Theobionine has a very lotter taste, which is only slowly developed. Its physiological action is similar to that of effects but more powerful. In large doses it produces well-defined poisonous effects. It is eliminated by the kidneys, and can be detected in the nume.

Theobremuse dissolves in 1800 parts of ne-cold or 148 of boiling water. In cold alcobol also it is only very slightly soluble (1 in 4280), and requires failly 400 parts at the boiling-point, but dissolves far more easily in 80 per cent spirit. If requires 1700 parts of cold or 600 of boiling other for solution, dissolves in 100 parts or boiling obleroform, is soluble in amylic alcohol, dissolves slightly in bearene, and is insoluble in petroleum spirit.

Theobromine dissolves in acids, and is precipitated from the solution by alkalies, but is soluble in excess of ammonia or fixed alkalies. It is wholly extracted from its solution in caustic soda by agitation with chloroform.

Theobromus is a weak base, its saits being readily decomposed by water with separation of the alkalod (compano Caffone, page 482). The hydrochlorids, BHG1+H₂O, and nitrate, BHNO₂, lose all their and at 100° B,H₂CH(2.₁ + 2H₂O orystallises in oblique prisms, which efforeses in the air and become anhydrous at 100° BHAUCI, forms triffs of yellow needles

An aqueous solution of the obtomine forms with mercune chloride a white crystalline precipitate, sparingly soluble in water and alcohol.

One of the most definite and machible compounds of the obscines a bat with nature of allow. When a very dubte appears solution of the obscines nature of the three properties of the solution of the silver-white needles containing C_1H_2N/O_2AgNO_3 form after a short time. The compound is only spannigly solution in water, and may be dread without change at 100°. If a solution of the obscine of the solution of the obscine obscine of the obscine obscine of the obscine obsci Theobrothane reacts with alkales like a weak and and forms definite salts. Thus the sectam salt is obtained by abling theobronnes to soul-ipe until a portion remains undissolved after long standing, and evaponising the fillate under the an-pump. The product is destinate of cystalline structure, is extremely soluble in water, has a strong alkaline reaction, and absolue calbon discard from the an The barium salt, (G-LHAN), Da, a separates on adding theobronnes to baryta-waten as a mass of microscopic mealing, and is obtainable as a snow-white field of silky necdicely by slowly cooling its solution in hot water. If the solution be rapidly cooled, it solutions to a significant could be solved.

Theobromme yields no product similar to caffeidine when boiled with concentrated baryta-water or caustic alkalies By such treatment, as also when heated with hydrochlone and under pressure to 240°, theobromme gives the same moducks as caffeine (uage 478).

The best preepition of theohouses as a solution of solution phosphomegasted (1992 186), which should be added to a solution strongly accidated with sulphane on artre soil. The yellow parespitate should be mixed with sulphane on artre soil. The yellow acceptate should be mixed with solution can be about or magnuss, draef, and the mixture exhausted with chiloroform, which dissolves the theobromany.

When theobrowme is heated with dilate subjutue each and lead duxule, exhon duxade is evolved. When once stated, the reaction continues without further application of heat, and it excess of the exclaining agent and too long heating be avoided the fiftered liquid is colouless, but ordere ammona on facations with a caustic alkah, separates subjutur from subjuturetted hydrogen, colours the skim purple-red, and immediately times blue when intacted with a moderate quantity of magnessa. Excess of magnesia destroys the colour, which may be restored by cautious adultion of subjuture need, colour, which may be restored by cautious adultion of subjuture need.

By oxidation with chrome and mixture, the obsoroum yields earlied outside, methylamme, and methyl-para bear o act, Q.H.(CH,N), Q.¹ Aqueous chloune converts it into methyl-ures, CHL(CH,N), and methyl-alloxan, Q.H.(CH,N), Q., while treatment with hydrochlore scut and potassium chlorate oxidases it to dimethyl-alloxanian, Q.H.(CH,N), X.Q., The obsomine gives with oxidiang agents and ammonia the same colour-reactions which characterises of flue to face 480)

ISOLATION AND DETERMINATION OF THEOBROMINE,

Theobiomine may be isolated by much the same methods as those

Methyl-parabanic acid is easily soluble in hot wats, from which it crystallises in transparent prisms, melting at 148° Waimed with aminoma and calcium chlorida, it gives a precipitate of calcium oxalate (compare Choles trophane, page 431). used for the determination of caffaine, having regard to the ful less ready solubility of the forms alkinded in water, alcohol, and other solvenia. As in the case of caffaine, the methods used by observers who have secorded high yields of theophonium are more trustworthy than those of eleminate who have succeeded in isolating commentatively small proportions.

For the preparation of theohomme, E So hm 1 dt (Archve der Pharmacie, coxxx 656) mixes commercial cocca (freed as far as possible from fat by pressure) with half its weight of frealityslaked lime, and extracts the mixture with boiling alcohol of 80 per cent (by volume) On cooling the alcohole extract, theobronine separates out, and on recrystallisation from hot alcohol is obtaned as a white, circulain analydous product.

Before extracting theobromme it is preferable to get rid of the fat by exhausting the inely-divided cocoa with petroleum spirit. The residue is made into a paste with water and ginited magnesia, dried at 100°, and exhausted with spirit of 80 rer cent

Another method of extracting the theolonomic from cocon is to exhaust the substance with water or dilute alcolod, pecupitate the solution with acetate of lead, separate the lead from the filtered solution by sulphuretted hydrogen, ovaponate the filterate to dynamic and extract the theolonomic from the readuc by boiling chloridom.

Caffeine may be separated from theohomine by treating the mixed alkaloids with cold benzene, in which theobronime is practically insoluble.

James Bell (Poods, 185) bolls 100 grams of the occas repeatedly with benzel, windot knostves fatty matters and onfilme.² The residue is mixed in a motter with 100 grams each of sand and calcuned imageness and sufficient water to form a paste, the product direct at 100°, and repeatedly boiled with strong alcohol Ins solution is filtered, dividueld, and the residual theohomine dred at 100° and weighted. It is freed from tances of fat and offence by teachment with hot bearner, and then tosted twee with

¹ By using a known volume of liquid and liftening off four fiftis or other known proportion, the tedions washing of the bulky lead quoenistic may be avoided. When once the alkaloid is in solution, the method recommended by the enthor for the determination of cafficine (page 490) is also applicable to thoshomine. The oblivoforms about he used was not support to the observations of the case of th

³ B.H. refers to the product, which was expectally yielded by Tunudal coses, as a "them bike alkaload," but as W or go an an and E Solvan of that base both proved the occurrence of coffeene in coool (Amadea, carra, 300) then o seems no doubt as to the nature of the substance observed by Bell Ho separated it from the fairly matter by boiling with water. The agreeous liquid was eraporated, and the alkaload punified by successive solution in water and banacaes.

a httle ree-cold water It is thus obtained white and perfectly pure, except for the presence of a trace of mineral matter 1 Bell found by this process the following proportions of alkaloid in cocca —

' Cocca	Theelaomine.	Theine like Alkaloid (Caffeine)
Guayaquil (nths),	0 54 per cent	Trace
Grenada (nibs),	0.91	Trace
Surinam (nibs),	0.78 ,,	0 02 per cent
Tunidad (nibs),	0.59 ,,	0 25 4
Trinidad (husks),	102 11	0.88 ,,
Trinidad (husks),		0.88 ,,

It is probable that Bell's issults are considerably below the tuth, since Pay on found 2 per cent, Mitselberlich, 15 per cent, Trojanowski, 12 to 46 per cent, while G. Wolfram found, max samples of dred coon-beams divested of their shells, from 134 to 166 per cent of theobrounae, with an everage of 156 per cent. The dred husks of the same beams contained from 042 to 111 per cent of theobrounae, with an everage of 156 per cent. We sign an I mound 047 per cent of caffining in the kernel and from 011 to 013 per cent in the shell of coons-beams.

G Wolfra m (Dwoif Polyt Jour, cxxx 240) has described the following method of determining theobronine ² If shelled cocco-beams are to be sandpred, they are ground up in a hot mortar to a thick peste a tegrammes of this mass or 30 grammes woight of chocolate is digested for some time in hot water, and the solution filtered. The filtrate is precipited with ammoniscal accists of lead, the solution filtered hot, and the precipitate washed with boiling water till the washings (ascidiated with nitice acid) cease to give a yellow precipitate with Schelbler's reagent (page 136). The filtrate is rondered slightly alkaline with softs, concentrated to slout 50 e., strongly acidiated with sulphure acid, and the lead sulphate separated by filtration. The filtrate is now tested with

² A similar method has been successfully employed by Mitscherlich for the isolation of theobronius from nime

This might readily be sumored by dissolving the alkaled in hot chloroform, and each treatment would obviate the necessity of treating the impute silk-sloid with water, which cannot be perfound without less Bell's process is neally inhancial with that personally described by Trops in wasky (Arbarre, [3] x 3, Jour Chine, See, x xxii 383, see, see, for the substitution of "bennol" for performent their, a change which suggests continued between the two ovlerate, and probably causes loss of thebrowners.

a largo excess of sodium phosphotungstate, (Scheibler's reagent) The ecagulation of the slimy, yellowish-white precipitate of theobromine phosphotungstate is facilitated by warming and stirring the mixture gently After standing several hours, the precipitate is filtered off and washed with dilute sulphuric acid (6 to 8 per cent. H.SO.) Wolfram then decomposes the precupitate by hot baryta-water, precipitates the filtrate with sulphune acid, removes the excess of the latter by barium carbonate, evaporates the filtered hourd, and weighs the residual theobromine, which is then ignited and any ash deducted. L Legler (Zestschr Anal, Chem, xxiii 89) dissolves the precipitate in caustic soda free from chlorides, nearly neutralises with sulphuric acid, evaporates to dryness with sand, and extracts the residue with amylic alcohol The solution is evaporated to dryness at 100°, the residue weighed, and the loss of weight on ignition regarded as theobromme. A preferable plan to either would be to mix the moist theobromine phosphotungstate with sodium carbonate, dry, and extract with boiling chloroform, which on evaporation would leave the theobromine in a pure state

Divertin Under this name a pieparation has been introduced into medicine having the constitution of a combination of sodium-theobronine and sodium salicylate, and the formula C.H.N.N.O.,C.H.(OH) COONa

Diurstin is colourless, odourless, sightly soluble in cold water, and meloible in chloroform or ether, but readily soluble in hot water or warm diute alcohol. The physiological action of diuretin is said to be guite distinct from that of the analogous compound of caffaine. It is stated to be much more readily absorbed than simple theoreomies, and to be devoid of any toxic properse, or of the peculiar excitant influence on the central nervous system exerted by refficience.

Owing to the high price of theobronine as compared with criffens, substitution of the former by the latter alkhold is possible, and hence G Vulpius (Jose Ohem Soc, lvin, 1475) has proceed the following method for the assay of duction—2 grammes weight of the sample is dissolved in 10 cc of water in a porcelain dush, the solution acidiated with hydrochloric acid, and then rendered family slikaline with ammonia. The inquid is kept for three lours at the ordinary temperature, and frequently stirred. The separated theobronine is then collected on a taref filter, the filtrate being used to transfer the last of the mother-luquor, and that theoforomine is then collected on a taref filter, the Gentle suction is used to remove the last of the mother-luquor, and that theoforomine is then washed twee with 10 cc of cold water, drad at 100°, and waighed By this method, Vulprus voi, III parts.

recovered from 41 to 411 per cent, of theobromme from pure duretin, 64 per cent, remaining in the filtrate and washings Making this allowance, the theologinus should not be less than 464 per cent, and that isolated should melt when carefully heated. be completely volatile, and dissolve readily in caustic soda solution From the filtrate from the theobiomine, the salicyhe acid can be isolated by acidulating with hydrochloric acid and agitating with chloroform The separated chloroform is washed with water to remove numeral acid, a little water and a drop of phenolphthalein solution added, and the bound then intrated with deconormal caustic Eich cc of walkali required for neutralisation represents 0 0138 gramme of salieving and Diuretin should contain 384 per cent of salicylic acid. The titration completed, the chloroform may be senarated and evaporated, when the residue will represent the 6.5 per cent of theobromine not previously separated, together with any caffeine the preparation may have contained. To prove the absence of caffeine in diuretin, Vulnius recommends that I gramme of the sample should be dissolved in 5 cc, of water, and the solution neutralised with hydrochloric acid, when the theobromine will form a milky precipitate readily soluble in soda solution. If the mixture he shaken with its own measure of chloroform, not more than 0 005 mamme of residue should remain on evaporating the separated chloroform

THEOPETRIANS, C.H.M.O.g. a base existing in minute quantity in tea, is isomeric with the obrowine and paraxapthine (occurring in himman unio). According to A.K. ose el' (Berichta, Xxi 2164, Pharm Jour, [3], xxi 41, Jour Othem Soc., iv. xxi 2164, Pharm Jour, [3], xxi 41, Jour Othem Soc., iv. xxi 2116, theophylime crystallises with laque, which it loses at 110° It mells at 264°. It is easily soluble in warm water, but spaningly in cold alcohol, and is extremely soluble in very dilute ammonia. It forms a crystalline hydrochloride, nitrote, diloro-

For its solution, K os sel extracts tan leaves with alcohol and evaporates that tentien to a synty, when need of the calleme expetituless cut on cooling. The filtrate is adiated with water, acaditated with sulphure acid, filtered after a considerable time, made alliants with summary and precipitated with intrate of sulver. After standing trenty for bours the precipitate is filtered attention of and wanted with inties and, on cooling this planul, the sulver interest compounds of advance and hypercentains (surema) expetitions out: The sent compounds of advance and hypercentains (surema) expetitions out: The sent soundainted with minus and and decomposed by sulphuretted hydrogen. On concentrating the filtrate, suremakes first cyraliness, and subsequently theophylitrates of the suremakes of the suremakes and being nearly insulations in the production of the production of the production is then superior the suremakes and being nearly insulational with one. The precapitation is then superior and being nearly insulational with one of the production of the produc

platinate, any-chlouda, and mercur-chlorida, and combanes with scale to form a readily soluble compound. When evaporated with chlorine-water, theophylline yields a scarlet readule, changed to violet on addition of animous. The situer-densities, $C_1H_1A_0N_1C_2$ is obtained as an anonphous presentate on adding uliver intraction an animous solution of theophylline. It crystallines from hot ammonia, and dissolves readily in intric send. The methyline derivative, $C_1H_2A_0N_1C_2$, proporate by heating the last statement with methyl points and methyl alcohol, mells at 229°, and as identical with a affains.

Tea.

The tes of commerce is the prepared leaf of Thea snessis (and Derhaps allied species), a shrub-like plant belonging to the gonus Camelita I to occurs native in the Himalayas and Assam, has long been cultivated in China and Japan, and is now raised largely in Rutish India Cevton Brant & 2⁸

It was formerly believed that green and black tass were the product of distinct plants, but it is now known that the difference is due to the method of preparation; black tea having undergone a certam amount of fermentation, whereas in green tea this change is carefully prevented. The leaves are gathered from the plants four times a year, and are distinguished according to their age. Each leaf is at first a "flowery Petces" leaf, which is the name applied to the leaf-bud. This becomes in succession "orange

1 French , le Thé German ; der Thee.

² The Report of H M. Customs for 1891 to 1892 states that the weight of tea imported from the pennesula of Hindostan showed a decrease of three million pounds, while that from Ception innensed by more than sixtem millions of pounds, exceeding for the first time that of China tas, which now forms only one fourth of our enters communities.

* "For Nach tras, the leaves are withered a little, relied to libeate the pures, let in talls for the proper state of Fernmentain, then usur-drast and sulpoted to a caseful firing in a furnace. For press tras, the fresh leaves are first withered in begs, and returned to the pans for a final abov rousting, with stiming, fresh and returned to the pans for a final abov rousting, with stiming, fresh to make however, and returned to the pans for a final abov rousting, with stiming, fresh of the pans for a final abov rousting, with stiming, for spit or man bounts, beginning at the temperature of 160° F., and falling to 150° F. the closes (A B Fraccott). The methods of preparing tea vay maternally in different countries. In India, the manufacturing processes are very much simplified, being related to five, instead of the twelver proctaced in that the leaves are not touched by the laboures, coursely in pricing. This is partially tree also of Japanese tas, whereas Climese to as manuplated dimental cantrely by hand, except when the feet are employed. A databled descomption of the method of proparing Japanese tea has been given by J. Takyams (Oken. Nices.). 1399)

Peton," "Peton," "Souchong 1st," "Souchong 2nd," "Congon," and finally "Bobas" ¹I mose access the leaves are classified emply as Peton, Souchong, and Bohea The first and second pucking of the season furnish the finest tess; but the quality of the protect depends on the age of the tace as well as the age of the leaf, the finest tess being produced from the young leaves of young plants, whilst old leaves, and the leaves of old wood, are deficient both in flavour and extract²

Besades the foregoing distinctions, based on the age of the leaf, there are other classifications based on the district of growth and the method of preparation. Thus among the chief commercial varieties of shade to are a Sasum, Caylon, Japan, Kassow, Moning, and Colong, and those of green tea, Gunpowder, Hyson, Young Hyson, Impensal, and Twansky Green tea has much declined in popularity of late years "Caper tea" is always more or less of a factions channels.

Vary few trustworthy complete analyses of tea have been pubhated; and, indeed, they have but a limited interest or practical value, since the tea is not consumed as a whole, but invariely, mitused, and the infusion contains the tea-constituents in very different proportions from those in which they exist in the lost An average of existen analyses of less made by Konir

showed -Moisture, 11 49 per cent , caffcine, 1 35 , albuminous

1 Pck-ho eignifies "white hairs," Sou-chong, "little plant," and Con-gou, "labour"

³ O. Kelluer (Lond Fession-Stat, 1988, 270, Jour Olem Soc, in 178, has published analyses of the leaves of the same to spiral dump or months (May to November) His figures show a decrease in the proportion of total larges, and showed entire dampesparance of annoth on togos in the following the leaves. The caffisms fell from 2 85 to 100 (satimated by evaporating the intension to dynams with magness, and extracting under sheele), and the tomation to dynams with magness, and extracting under sheele), and the summon for the size of the state of the size of the siz

The albummonis were detartuned by a modification of Statzer's process. The squeous decorbon of 2 grammes in 100 c of water was treated with 20 c c of a 10 per cent column of cuprac enliphate, and a finisted solution of caracter cold in used quentity as to leave a title copper in solution. The promised buyland fibract lapidity, and was froe from albummonds. The promised washed first with hot water and then with strong alcohol. The contained introgen was determined by garinos with cold-lines.

matters, 22.22, ethereal oil. 0.67, gum and dextin. 7.13, tannin, 1236, fat. wax, and chlorophyll, 362; other nitrogenfree matters, 16.75, woody fibre, 20.30, and ash, 5.11 per cent

J M Eder (Dingl. Point Jour. ccxxxx, 445, 526) gives the following as the average composition of tea.2

A	Soluble in mater-	Per cent	B Insoluble in reates	Per cent
	Moisture,	100	Chlorophyll,	1 S to 2 2
	Tannin,	. 100	Wax, .	0 2
	Gallic acid, oxalic	acid, and } 02	Rosin,	30
	quercetin,	. } "	Colouring matter,	. 18
	Boheic acid, .	01	Extractive matter, me	atly } 160
	Caffeine or thomo,		solubie in nitrie acid,	,
	Tea oil,	. 08	Cellulose,	. 200
	Albummous bodies	(probably)	Albuminous hodies, .	. 127
			Mineral matters	4.0
	Gummy substance and sugar,	s, dextrin, }s to 4		
	Mineral matters,	. 17		

ESSENTIAL OIL is determined by distilling a considerable quantity of tea. (200 grammes) with 1500 o c of water, and agitating the distillate with ether On distilling off the ether the tea oil remains Eden found 0 52 per cent. of oil in guinewder and 0.41 per cent, in pekoe bloom tea by this mocess Battershall employs 10 grammes of tea and saturates the distillate with calcium chloride before agritating with other. A good sample of black tea violded 0.87 per cent of volatile oil when examined by this method

Tes oil is a bright vellow liquid, which darkons and issimiles on exposure to the an fer a few days, and turns reddish brown with mitrio acid. Even on expesing the aqueous distillate from tes to the au for some time, it loses its alomatic edour, and little or no oil can then be senarated from it by ether. and even if the distillate be kept in closed vessels the alome is com lost. These facts explain the fact that tea leaves lose their bouquet by age or expesure.

Quercitrin and Quercitin, stated by Hlasiwetz to be mesent in tea, are described in Vol III, Part I page 341

Beheld Acid, C.H.,O., according to Rochleder (Annales, lvin 202). exists to the extent of 0 1 to 0 2 per cont in black tea. It is memoral by precunstating a boiling intusion of tea with accetate of lead, neutralising the filtered liquid with ammonia, susponding the washed precipitate in absolute alcohol, and decomposing it by sulphuretted hydrogen. The filtrate is evaporated to diviness in vacuo, and the residual bohere acid purified by resolution in water, &c It is a yellow resinous body, melting at 100° to a tenacious manner, and decomposed on exposure to air It is extremely soluble in water and alcohol, and giving a brown coloration but no precipitate with ferric chloride The salte are mostly insoluble and amorphous

² Edor's figures for mineral matters soluble in water are considerably lower than any other observer, and his proportion of insoluble matters is in excess and of soluble in definency of those usually recorded. His tannin, which was The following analyses by Y Korai (Bulletin, No 7, Imperial College of Agneulture, Japan) have a special value owing to the author's knowledge of tea manufacture. Special piecultions were taken in amphing the leaves to ensure strictly parallel specimens being taken. The figures refer to the moisture-free leaves in each case.—

			Unprepared Leaves	Green Tea,	Black Tea
Cullefne or theine,	_		8 30	8 20	8 80
Ether extract,			6 49	5 52	5 82
Hot-water extract, .			50 97	58 74	47 28
Tannin (as gallotannic acid)	,		12 91	10 64	4 80
Other nitrogen free extract,			27 86	81.43	85 89
Crude protein,			87 88	37 48	88 99
Crude fibre,			10 44	10 08	10 07
Aah,		•	4 07	4 02	4 93
Albuminoid nitrogen,			411	8 94	4 11
Caffeine nitrogen,			0.90	0.88	0.98
Amido-nitrogen,			0 91	1 18	1 16
Total nitrogen,			5 97	\$ 09	6 22

The proportion of ash found by Kozai is remarkably low, but it seems not impossible that this is characteristic of Japanese teas, since some analyses by J. Takayama (Chem Neus, 1 299) show the same peculiarity.

An analysis of the so-called "flower of tea," consisting of the hairs of the leaf-buds of the tea-plant, has been published by T B Groves (Year-Book Pharm, 1876, page 610)

James Bell (Foods, 1. 6) gives the following figures as illustrating the composition of fair representatives of black and green tess of commerce 1—

determined by precupations with cupue costate, is unascully low. Of the extract, from 15 to 16 per cent was precupable by strong alcohol. A nitrogen determination on the precupating gave a result corresponding to about 12 per cent. of alluminous matters, and the difficience was regarded as genmary sail scalences. The otherphyll, war, and resun were extracted by other from the unsoluble natter, after drying, and the resultant celluloses purified by treatment with intrus onel, petash, and alcohol.

¹ It is evident that in these analyses some constituent was determined by difference, but it is not stated which. Not does Bell state the method used for determining the tannin, the figures for which are highly improbable, while other of his descriptions are meanifeld or obscute.

								(Black)	Young Hyson (Green)
Moisture,			_			•		8 20	5 96
Caffeino, .								8 24	2 83
Albumin, insoluble, .								17 20	16 88
Albumin, soluble, .								70	80
Extractive by alcohol, c	nta	ining	nite	ogen	ous 1	natte	er,	0 79	7 05
Dextrin or gum,									60
Pectin and pectic acid,								2 00	8 22
Tanniu,								16 40	27 14
Chlorophyll and resm,								4 60	4 20
Cellulose aud insoluble	oto	ıring	mat	ter,			. 1	34 00	25 90
Ash,								6 27	6 07
								100 00	100 00

The following figures are given by J. P. Battershall (Food Adulteration, page 28) as the results of the analysis by American chemists of sumples representing 2414 packages of Indian tas, a class remarkable for their general staength, high quality, and freedom from adulteration.—

	Minimum	Maximum	Average.	
Moisture,	5 88 per cent.	6 82 per cent.	5 04 per cent	
Insoluble leaf,	47 12 ,,	55 87 ,,	5191 "	
Extract,	87 80 ,,	40 35 "	3881 ,,	
Tannin,	1304 "	18 87 ,,	15 92 ,,	
Caffeine on theme,	188 ,,	3 24 "	274 ,,	
Ash -Tutal,	505 ,,	6 02 ,,	561 ,,	
Soluble in water,	3 12 ,	4 28 ,,	3 52 ,,	
Insoluble in acid,	0 12 ,,	0.80 "	0 18 ,,	

The following figures, obtained by C M Caines in the author's laboratory, are interesting as indicating the character of the first parcel of Natal tea ever imported into England 1—Moisture, 8 36

¹ Natal tea must not be mustaken for the so-called "Capo tea" and "Bush tea," consisting of the dired leaves and twigs of contain species of Coglopus According to H. G. Gleenish (Pharm Jour, [3], in 549, 569, 882). Cape tea is destrute of cassens or other alkaloid, but contains a

per cent , insoluble matter, 51 96 , hot-water extract (complete), 39 68 , tainin by PhÅ₂ a 33 , tainin by CuÅ₂ 8 50 , caffeine by PhÅ₂ and chloroform, 2 85 ; total ash, 6 14 , soluble ash, 3 56 , alkalmity (K₂O) of soluble ash, 1 15 per cent,

The Mossture contained in commercial tea varies within some what wise limits (4 ± 10 18 per cents), but the range is far less when tess of the same cless are compared. Thus G W N₁g, nor (Pham Jon. [3], vi 381) found that hyson and gunprowlars, both of which are highly-drued teas, contained the smallest proportions of mossiture (4 8 to 6 55 per cent), and, after drying at 100°, absorbed from 604 to 6 98 per cent of water on exposure to are Congost teas showed in their original condition an average of 8 50 per cent of mossture, and never wholly reguined their original weight on exposure to air after drying at 100°. The average proportion of mossture in commercial tea is about 7.7 per cent, and the mean Launce between T and 9 per cent.

CAFFEINE OF THEINE The proportion of alkaloid present in tea varies considerably, the general range being from 3 0 to 4 0 per cent. ; but the experiments of Paul and Cowuley (page 492) show that in Indian and Ceylon tea the proportion is more frequently above 4 per cent than below that figure, and in a special sample of Himalayan tea. Zoller found as much as 4'94 per cent of caffeine, in addition to a small proportion of what was apparently theobromine Unfortunately, by far the greater number of published determinetions of caffeine are quite unreliable (see page 484), and, indeed, the low figures recorded suffice to indicate their inaccuracy, and hence any deductions as to the relation of the quality of tea to the proportion of alkaloid present must be received with great caution. The proportion of caffeine is not generally considered to have any direct relation to the commercial value of the tea, and the tea-taster takes no cognisance of it. The results of J F Geisler (page 506) tend to show that the proportion of caffeine which passes into the infusion has a relation to the quality of the tea, the superior qualities giving up their alkaloid to water more perfectly than the inferior, but as the whole of Geisler's figures for caffeine (1 15 to 3 50 per cent) are probably below the truth, too

glicossidal body called cyclopin, $\Omega_{\rm eff} M_{\rm eff} \Omega_{\rm o}$, smulter to unchomanovatama each, and yelding, when bolled with dilate end, glicoso, and a unbrance of the formula $G_{\rm eff} H_{\rm eff} \Omega_{\rm o}$ decoly resembling nuchona-normarial General absolutes are being a cycloping and the cycloping and the preventagl described by Δ if Ω therein (Ω few, viz. 18), and likegroundly described by Δ if Ω therein (Ω few, viz. 18), and likeproduct of that body. One to yielded the author 50 per cent. of actuer, and on quiction left 37 per cent. of an each contains managements. much stress must not bo land on this conclusion, I and the sime remarks applicable to the proposition of P D via k vi to b, that the higher the proportion of alkaloud bears to that of the samma and fermantiation-products, the more valuable is the tea. This varied from 16 0.84 to 24.5, 75 5, the percentage of alkaloud in the host belf remarks from 24 to 3.45 for example.

CRITGORISTIA. When either green or black tes is bolled with alcohol or chlordorm a solution of a more or less grass-gron colour is obtained, owing to the extraction of chlorophyll. E. B. Kenrick states that chesp black tess yield less chlorophyll than the better linds, and believes that a distinction of practical value might broably be based on a colormistic determination.

Extract By the ten "extract," when used in reference to ke analysis, is understood the sum of the soluble matters extracted from the leaf by bothing water. It includes cafferine, tannin, albuminous matters, gum, dextrin, colouring matter, mineral matter, &c, beades other less important constituents, such as gallic seed, behave and, coxide seed, and querosin, which substances are unseent in commarative small cannity, if at all

The proportion of extractive malite yielded necessarily vanes with the method used to exhaust the tea, and is, of course, higher when the ten is powdered and the treatment with water long contanced and carried to an extreme than when the whole leaves are used and the tea simply influeed in boiling water. The latter method commends itself when the object is to study the character of the infusion likely to be yielded in practice, while the former plan gives more information when the objection is the detection of scallers too.

An interesting comparison of the results of the two methods has been made by J. F. Ge 1s1 e.g., who has published an extensive series of analyses of teas obtained direct from American importees and wholeash houses (American Grocer, Oct 23, 1884; Analysix 20), Proisout's Organic Analysis, page 505 et 3eq). The following table by Gesler shows the proportions of extacts, tannin, caffeine, and ash which passed into solution when various representative commercial leas were infused under precisely the same conditions by pouring ou the leaves 100 times their weight of boiling distilled water, and allowing the higher in 'dian' for ten minutes. The ratio which this dissolved constituent bore to the total is also given by

¹ In a private communication to the author, Mr Gesler states that the caffeine was detainmed by mixing the concentrated docection with magnesia and said, and exhausting the dry mixture with chlorofoim (compare page 486).

	Wholesslo	Ext	raot	Tannin 2		Caf- feine	Ash	
Kind of Tea.	Price per lb in Cents	Infu- sion	Ratio to Total	Infu	Estio to Total	Infu slou	Infu ston	Ratio to Total
Fine Ceylon Pekes trps, 1 Assam, Assam, Finest Mayune Gunpowder, Common Movuso Gun	234 231 76	33 25 29 15 28 67 37 32	76 6 78 6 72 0 78 2	17 19 11 48 9 50 16 79 9 20	70 3 00 8 58 1 87 8 77 7	2 44 8 80 2 75 2 05 1 67	8 11 8 80 4 40 4 00 4 00	91 0 70 0 79 5 55 8 60 1
powdes, Japan Basket fired, Japan Pan fired, Japan Pan fired, Choic est Formosa Oolong, Choicest Formosa Oolong, Superior Formosa Oolong, Superior Formosa Oolong,	65 58 30 24 21 45	28 07 81 75 84 37 83 02 33 30 20 00 27 40 24 50 24 26	79 4 75 8 79 6 75 9 73 7 08 0 60 9 60 5	11 28 19 41 12 01 13 76 9 03 10 12 7 53 5-46	74 5 94 4 75 6 68 5 50 0 50 0 55 6 41 7	2 17 2 07 2 50 2 42 2 12 1 92 1 70 2 87	4 27 8 67 4 00 3 07 8 60 8 72 8 25 4 18	80 8 03 6 71 3 00 5 02 3 68 5 58 0 78 7
Superior Moning Congou, Modum Moning Congou, Good Common Estsow Con- gon,	27 103 174	21 55 21 02 28 25	68 6 64 1	4 41 5 55 4 05	\$2.0 45.2 88.5	2-77 2-83 2-35	8 70 8 22 3 30	03 5 58 8
Common Moning Congou,	151	19 50	72 2	4 50	52 9	1 95	2 88	40 8

- 1 Jour American Chem Soc, aitt, No 8
- 2 The determinations of tannin were made by the Löwenthal method, except in a few natures in which the course seet its method was employed.
- 3 This sample is considered by Goisler to have been adulterated, though its appearance did not indicate any admixture with exhausted longe—(frivate Communication to Author)

A comparison of these figures shows that, as a rule, the fine near leas yield to hot water larger proportions of extract, diffens, and sah than the inferior qualities. On an average, the sah of the extract exceeds by 0.62 per cent the "soluble sah" obtained by teating the ash of the entire tea with water. The proportion of tanimn rises and falls with that of the extract, and the ratio of which the dissolved extract and tanimn bear to the total has a notable relation to the price of the test.

By the same method of 10 minutes infusion in boiling-hot wates, E B Kenrick (Bulleton No 24, Laboratory of Inland Revenue Department, Canada) obtained the following average results from commercial samples of tea —

Description of Toss	No. of Samples	Aqueous Extract	Tannın Dissolvod	Cafferno Dissolved	Ratio of Aq Extract to Tannin
Congou,	10	23 37	5 18	2 65	4 51
Assam,	8	88 53	7 40	8 98	3 81
Coylon,	2	97 45	7 85	2 68	8 50
Unclassed Black,	13	23 76	5 40	2 82	4 40
Japan,	18	80 07	9 88	2 45	8 20
Gunpowder,	2	28 56	8 90	2 39	3 67
Young Hyson,	5	84 22	10 98	2 52	8 12

From these figures it appears that congon teas yield less extract than green and Japan tess, while Assum and Ceylon teas yield intermediate results. Not only do the Japan and green teas yield more soluble tannin than the black, but the proportion of tannin to the whole extract is greater in the former kinds. On the other hand, the black teas appear to yield more soluble caffeine than the Janan and green teas.

The following figures by Geisler show the influence of the time allowed for influidin upon the proportion of the constitution dissolved, and the difference in the result caused by substituting Now York water (Croton River, of 49 6 degrees hardness per 100,000 for distilled water. In each case the tes used was the finest Formose Oclong, and it was infused in 100 parts of boling water—

		Distilled	Croton Water			
	8 min	5 min	10 min	1 hom	5 min	10 min
Total extract,	25 97	28 87	30 87	88 75	27 47	90 25
Ash,	3 72	8 80	4 17	4 33	3 62	4 13
Extract menus ash,	22 25	24 60	28 70	20 42	23 85	26 12
Tannin,	9 75	11 23	13 46	14 94	10 18	10 60
Caffeine,	1.05	2 65	275	2.85	2 02	2.82
Alkalinity of infusion ash ($=$ \mathbb{K}_2 0),	1 03	1 08	1 22	1 28	108	1 15

From these results it appears that infusion in distilled water for a minute is in smifitnent, but in 5 minutes practically as good a result in distanced as na longer time, without so much astringed matter boing extraded. When Crotor water is used; 10 minutes gives a maternally better result, so far as suffisine and extinct are conserned, while the proportion of stamm is not increased in the same proportion. In all these experiments the voltation of the form of consideration, though it is to thus constituent that the flavour and aroun of the ten as dua, and on these characters the commencative matter impart astungency, strength, and body to the minute of Caffann, busing almost tasteless, is not taken mot account by testastics, though physiologically it is the most miportant constituent of test.

In tasting tea, it is usual to infuse the weight of a sixpenny piece (43 grains) of the sample in 3½ fluid ounces of boiling water, and to pour off the infusion after standing from 3 to 5 minutes, according to the practice of the taster The infusion is not swallowed, and, of course, no sugar or milk is added. In the process of manufacture, the different sized leaves are senirated by sifting, and thus broken leaves and dust are obtained, which though vielding a strong infusion, will be sold at a lower rate Broken or powdered ten loses its aroma more sandly than wholeleaf toa Hence, in judging of the commercial value of a tea, the appearance of the leaf and extent to which it is damaged are taken into account as well as the characters of the infusion. The infusion is judged by its strongth or astringency, its flavour, its colour, and its odour The strength and flavour are dependent on the age, and consequently the size of the leaf, and the time the tea has been kept since its manufacture. A chemical analysis will indicate the strength, but not the flavour of the infusion, and hence is of little use in the valuation of high-mixed teas, but as in medium and low-priced teas the strength is of as great or more importance than the flavour, a chemical analysis will in such cases. go far to indicate the commercial value of the tea. The common formed of a tea by a professional taster is somotimes very different from that to which a chemical examination would lead 1

It is comparatively unusual for unmixed tea of any kind to be sold retail. Blending of several kinds is very generally practised, and when conducted judiciously materially improves the character of the tea.

In 1874, the author submitted to two teatesters of considerable experience a series of samples which he had specially prepared to test their ability to recognize adulterations of tea by the tasts.

The following were the opinions expressed.

Nature of Sample	A's Opinion	B's Opinion
No 1 70 per cent of No 2 and 30 per cent exhausted and redried leaves.	Tasted "washed-out," no doubt from presence of exhausted leaves	Very poor, contained many exhausted leaves, ranked Afth
No 2 Genuine black tea of fair quality	Genuine	Passed pure , ranked fast
No 3 No 2 somewhat crushed	hixed with exhausted leaves	Would have been the bost, but lacks strongth, and is thorefore suggestive of exhausted leaves Ranked therefore
No 4 80 per cent of No 2 and 30 per cent of ex hausted leaves, to which a little Na ₂ CO ₂ was added while redrying	Genuine, better ton than No 3	
No 5 80 per cent of No 2, 20 per cent of exhausted leaves, and a little cuto chu	A washed out to a to which some astringent matter had been added	Passed pure, and ranked second

Adulterations of Tea

Before the passing of the Adultention of Food Act of 1872, which was subject to adulterations of the grossest kind, a most of which were procised prior to importation. By the Sale of Food and Drugs Act of 1875, provision was made for the examination of ten by the Custom House, and the exportation or destruction of vary bad parcals ² Hence the ten now sold in the United Kingdom is restricted adulterated in the gross manner which was formely common ²

The adulterants of tea may be conveniently arranged under the following four heads — I Mineral additions used for increasing weight or bulk, such as sand, magnetic iron one, bruse filings, 2 Organic abiditions used for increasing weight or bulk; such as previously infused leaves, and leaves othis than those of the tea plant, as slow, dider, willow, &c 3. Adulterants used for imparting fastitions strength, by increasing the astringency or deepring the colour of the infusion, as catechie, softime orborate, but 4 Feengs and colouing materials, as steatite, prussian blue, indige, turners, crashite, &c

The practice of facing tea, formerly very common, is now confined to certain kinds of green tea, especially gunpowder, and the

1 By section 5 of 11 Googs I cap 30, the adulteration of tea by terna, appoints (catedun), learned other than leaves of tea, or any other ingredients whatever, was punshable by festerine and a fine of £100. By section 11 of 4 George III, and 14, a penalty of £10 was imposed for the sale of every pound of tea which was mixed, coloured, stuned, or dyed with turns appoints, sugar, molasses, also, records or with any other ingredients or materials whatesome.

*On May 9, 1891, W. Go b d on S am o s, the church chemats in the Construction House Laboratory, neported that 457 amaples had been analyzed during the year 1890, viz. —85 samples green fixed 16 black dust tes, and 25 samples of affungs Of these, 885 samples were found on analyzes to be satisfactory. Of the creamang 68 samples, representing 50 ploakages of southful and unsound tess, 1 sample, 1 sepresenting 50 ploakages, were restricted to be expention, owing to the presence of exhausted leaves, damage, or other causes within the Act, 8 starting the contract of the

⁵ This statement does not apply to all countries. As recently as 1883, We on da and Wiologors ki doscribed various adulterations they had met with in teasold in Warsaw Bukowski and Aleksand 10 w in the same year found as much as 40 pu cent of ash in tea, and a considerable proportion of brass filings in one sample.

mmeral additions for mereasing weight or bulk no longer include (so far as the United Kingdom is concerned) considerable proportions of magnetic from one, &c., as was formerly the case

For the detection of mineral adultments, and to obtain certain other knowledge, the tea should be ignited, and the proportions of ash soluble in water and acid determined. In practice this is best effected by igniting 5 giammes of the tea, in its ordinary commercial condition, in platinum, at as low a temperature as possible. When the carbon is burnt off, the ash will have a distinct green colour, owing to the formation of manganate The ash is weighed and boiled with water, the solution filtered, and the residue washed, ignited, moistened with ammonium carbonate, very gently ignited, and weighed. The difference between the weight now found and that of the total ash gives that of the ash soluble in water 1 The insoluble ash is next boiled with strong hydrochloric acid, the solution diluted with hot water, filtered, and the insoluble residue washed, ignited, and weighed. It consists of extrineous siliceous matter, such as sand, fragments of quartz, &c, and insoluble silicates, such as steatite from the facing of gunnowder tea If titanic non sand be present, some of it will almost certainly remain undissolved, and present the appearance of jet-black magnetic particles 2

The solution of the ash soluble in water should be titrated with

¹ If preferred, the weight of the soluble ash can be ascertained directly, by evaporating the solution and weighing the residue.

2 The adulteration of tea with magnetic matter, formerly (in the experience of the author) very common, is now apparently nearly obsolute, a clear proof that the mineral admixtures were not due to accidental causes Magnetic matter is best detected by reducing 10 grammos of the tea to powdor and sproading it m a thin layer on a sheet of smooth paper. A magnet or electro-magnet is then applied to the under-side of the paper and moved laterally, with its poles in contact with the paper Any magnetic matter may thus be readily drawn out and separated from the tea. It is next boiled with water for a few minutes to detach adherent organic particles, and the water decented. The residue is thon dried and weighed, and examined under the microscope as an epaque object. If it consists of magnetic oxide or titanate of non, crystalline facets will probably be apparent, the bulk of the object having a jet black colour Metallic iron would be distinguished from the foregoing feringinous minerals by its colubility in moderately strong nitric acid (sp. gr. 12) with evolution of red fumes, and by its precipitating metallic copper from a warm and slightly acidulated solution of cumio sulphate

The weighting of the matter actually extracted by a magnet us fat more satisfactory than the estimation of the tree nectually amounts to about 3 per entantially contains a small proportion of row, but it only amounts to about 3 per cent of the weight of the six, or about 0.18 per cent of the entire tax Of course the tree in this form is not affected by a magnet, the use of which has the

methyl-orange or litmus and standard acid, the volume used being calculated to its equivalent of potassium oxide (1 cc of $\frac{N}{10}$ acid = 0 00471 gramme of K_0 O)

The analyses of a very large number of tess show that the proportion of soluble sats and its alkalanity vary with the age of the leaves, the figures yielded being highest with young leaves and tess of high quality. The total sat of absolutely pure tex result, if ever, exceeds 6 per cent, but some heence must be allowed in dealing with commercial samples. Hence in 1874, the Society of Public Analysts suggested 8 per cent as the maximum limit of Public Analysts suggested 8 per cent as the maximum limit of total sat allowable in the of which not less than 3 per cent, should be soluble in water. These figures refer to ten previously tirred at 100°, and as the proportion of water usually the between 3 and 8 per cent, the corresponding limits for tea in its criticary commercial condition will be 7 40 and 27 Type cent respectively.

Somewhat more recently (1875), G W W 1g is 1 (Pharm. Jour., [3], v. 262, 381) obtained the following average issuing the same state of the same state of the same state of the same state. The samples of commercial tea taken from the original cheets. The samples enbraced forty-one of ordinary character, eighteen special teas of high prace, and time samples of caper. Wigner regarded and described these last as "genuine," and they were clearly free from any large proportion of mineral adulterants, but the author strongly questions whether any specimen whatever of caper ten really describe the secreption of "genuine"

				- 1	Results of Analysis of Ash					
					Total	Siliceons Matter	Soluble in Water	Alkalinity as K ₂ 0		
Samples on Come	erna	Stat	o					-		
Maximum,					7 02	1 67	8 88	1 06		
Minimum,				. [5 17	0.01	2 64	1.08		
Average,					5 78	0 46	8 15	1 45		
Samples ofter år	nng a	100	c-	Ì]		j		
Maximum,					7 42	176	4 16	2 11		
Minimum,				Ì	5 57	0.04	2.04	1-26		
Avernge, .					6 33	0.50	8 45	1 54		

advantage of extracting the iron in the form in which it actually exists, and production in court if necessary.

In 1873 and 1874 the author floquently found from 5 to 8 per cent of magnetic matter in caper tea, and at that time the use of the magnet for its detection was well known to, and habitually practised by, the trade.

The ash of these sixty-seven samples of tea had the following average composition.

		Including Silics, &c	Exclusive of Siltes, Ac				
Silicons matter, Soluble in soid, Soluble in water,	•	:	:	:	:	7 96 per cent 37 54 ,, 54 50 ,,	per cent 40 79 ,, 50 21 ,,
Alkalinity of solubl	io asi	1, 81	K ₂	0,	 	100 00 per cont 25 00 per cent	100 00 per cent 27 26 per cent

J am e B B 11 (Foods, 1 29, 31) has published figures agreeing with those of Wignes. The proportion of soluble sak in genuine tess analysed by Bell ranged from 9 8 to 4 2 per cent (calculated on the moisture-free test), the proportion being usually believes 2 and 3 3 per cent. In one instance only did the soluble sak field below 3 per cent, and in that case the deficiency was very triffing, the proportion being 2 97 per cent. The alkalmity of the soluble sak of the tess excusioned by Bell ranged from 130 to 191 per cent of K_0O . In only one case did the total sah reach 8 per cent, while the innolluble subscore matter exceeded 1 per cost in a few matances only. Bell's results are fairly in accordance with the wine experience of the author (see Gham News, Xxx, 167, 189, 221).\(^1\)

¹ The following results of partial analyses of average samples of commercial black teas, as ordinarily imported, here been communicated to the author by M J Sheridan, Assistant Chemist in H.M Customs Labotatory The figures refet to the undired tea —

	1		Ash		Extract, on
Description of Tea		Total	Soluble in Water	Silicons Matter	Whole Leaves
Indian — Orange Pekoe, Asam Pekoe, Souchong, Pekoe Souchong,		5 40 6 10 5 70 5 76	8 20 3 80 8 15 8 25	0 45 0 90 0 80 0 70	40 49 30 82 80 44 86 78
OEYLON — Broken Orange Pekee, Ceylon Pekee, Soucheng,		5 50 5 40 5 00	8 20 2 85 8 40	0 20 0 25 0 80	42 90 38 24 37 98
JAPAN Siftings,	.]	6 12	3 15	0.05	29.60
JAVA Congou, Congou,	:	5 60 7 66	8 05 8 75	0.50 1.05	84 ¹ 60 80 72
CHIMA:— Kaisow Congou, Common Congou, Souchong, Octong, Flowery Pekoo,		5 70 5 85 5 80 5 85 5 45	8 26 2 16 8 06 8 20 3 06	0 50 1 00 0 65 0 70 0 65	82 95 81 71 83 57 84 10 85 70
PORT NATAL Congou,		5 65	8 10	0 45	84 80

In ortan cases a high soluble ash does not indicate a high quality of tea. This happens when the sah contains a notable proportion of sodium chlorde, owing to the tea having been damaged by sea-water and rodned. The saft of pure tea contains only a trifling proportion of sodium, less than 2 per cent, and the chlorine does not exceed 11 per cent, equivalent to about 18 do sodium chlorde, lepresenting 0 108 per cent of the weight of the tea Wigner (Pharm. Jour., [3], 71. 403) found 3 08 per cent, of sodium chlorde in tea which had been a fortunght under sea-water and completely soaked, and 0 17 and 0 23 in samples which had been a fortunght with the sample which had been a fortunght with the same of the sample which had been a fortunght was samples which had been a feathly mostened.

Prenously utilized or alhausted leaves are among the adultariations of tea most difficult to detect, especially when present only in moderate proportion. The sophistication of tes in this manner was formerly extensively practised in England, the exhausted leaves being frested with gum or other matters, and rolled and

redried so as to resemble genuine tea 1

The treatment of tea with hot water necessarily results in the removal of eartman of the sah-constituents, especially the potentian salts of organic acids. Hence the exhausted leaves will contain a smaller proportion of total sah, and especially of ash soluble in water. The extent of the change produced by infusion will, of course, depend on the perfection of the exhaustion. The sutting found in a mixture of infused leaves from vaccous teas 430 ns to Ball (Analysis and Asisteration of Foods, 129) gives the following figures obtained by the analyses of the sah of its eleaves which had been infused in the ordinary way for domestic use, and afterwards reduced at 100° —

	Ash of Sample							
Description of Tea.	Total	Sinceons Matters	Soluble in Water	Alkalmity, as Eg0				
Congou, .	8 92	0 41	0.54	0 11				
Moning,	4.63	0.96	0.86	0.28				
Orange Pelsoe, .	8 77	0 57	0.68	018				
Hyson, .	5 56	140	0.76	0.91				
Souchong,	4 12	070	0.81	0 19				

Though less extensively carried on than formerly, the practice of redrying univesal tea leaves is not obsolete. The infused tea-leaves from the various bread and kindled chops, now so numerous in London, are regularly VOL. III. PART II.

The total ash of the foregoing samples averages 4:38, and the soluble ash 0.73 per cent

Exhausted tea-leaves are also indicated by the deficient extract (and consequently high mesoluble maters) and low proposition of tamin. As already stated, the yield of extract depends materally on the condition of the tea, more complete extraction of the soluble matters being effected when the powdered tea is used than when the exhaustion is effected on the leaves in their commercial condition. For the purpose of detecting adulteration, the powdered tea should always be used, or the results will not be fairly compatable

The defenimination of the total extract and insoluble matter of the is best effected by boding 2 gammes of the less in a state of power with 100 c c of water for one how. The liquid is hittened while hot, the resultee bodied again with 30 c c of water, and the process repeated is long as colouring matter continues to be extincted, the liquid being powered through the filter previously used. After cooling, the descention is made up to 250 c, or often convenient measure, and an aliquid part (one-fifth) evaporated to dryness for the determination of the extract The fifter and its confess should be dried at 100°, and the misoluble matter detached and weighed. Very constant results are thus obtainable

The minimum proposition of extinct yielded by genuine teachanisted in a state of powder was fixed by the Sucasy of Public Analysis in 1874 at 30 per cent. Assuming the presence of 7.5 per cent of mostaire, this leaves 62 per cent for the maximum proposition of modulle matter. This figure covers almost all legitimate viruntions in tea, and is considerably in excess of the propertion yielded by green tes, the modulible matter from which averages 42 per cent, while in black tess the average is only about 50 per begitt up and reduced, and the leaves of the tea mutuch by textistins are

systematically preserved and sold for the same purpose.

1 J M Eden obtained the following figures by the analysis of tons adultrated with exhausted leaves purchased in small shops in the suburbs of Vienna—

		Tannin (by CuX ₂)	Extract	Total Ash	Soluble Ash
Russian ten,	•	6 00	18 4	4 76	0.85
Bloom tes,		4 91	15 8	8 84	0.54
Bloom tea,	•	5 18	14 6	4 51	0 90

² The decection of some teas filters very slowly, and it is necessary to strain the liquid through fine muslin instead of filtering it through paper.

cont In the case of obl-leaf Congou teas containing much stalk, and which have been stored for some time, the extract may occasionally fall to 30 per cent, corresponding to 63½ per cent of insoluble matter. In udging a ten by the proportion of extract or insoluble matter, it is vay desimble, when possible, to take into account the character of the sample. Thus young leaves (which are to some extent indicated by their sizely yield a notably higher extract than fully grown or old leaves, or specimens containing a considerable two proportion of state.

G W Wigner has recorded the proportions of extract yielded by a sample of team powder when boiled with different quantities of water. In each case the tea was boiled with the water under a reflux condenser for one hour, the decortion cooled, filtered, and evaporated to dryness.

Even after four boilings with 20 perts of water, the tea was not completely exhausted. Hence Wigner preferred to determine the extract by boiling the powdered tea once, for one hour, with 100 parts of water, instead of repeatedly exhausting with smaller quantities. Operating in this manner he obtained proportions of extact ranging from 26 15 to 44 85 per cent, the average being 35 70 per cent, containing 46 36 at al.

The deta manufact of tamass in tea affords valuable information respecting the probable presence of previously infused leaves or extraneous tamain matters, such as calculum. This is best effected in the aqueous decoction obtained by exhausting the sample with boling water, as required for the determination of the extract

The tannu may be estimated by H. R. Procter's modification of Lowenthal's process, as described in Vol III Past L page 110 A volume of the above desoction, corresponding to 04 gramme of tea, may be taken for the original tirtation with permanate, and of the desoction deprived of tannia a volume correspondi-

¹ The sals of the soluble extract of tea always exceeds by a considerable amount the proportion of tea sals soluble in wates, doubtless owing to the presence in ten of soluble salts of calcium and magnesims with organic solds, which salts on ignition are converted into calcium carbonate and magnesis, and thus become usefulbe in wates, ing to 0480 gamme of tea. The tamin of tea is stated by some chemists to be gallotamine and, and by others to be identical with that of oak-bark. The reduction-equivalent of the latter is almost identical with that of crystallised oxahe and, so that the weight of this substance corresponding to the volume of permanganate decolorised gives without calculation that of the tamin present.

The process of fermentation to which black ten his been subjusted undoubtedly eases modification of the tanum, with formation of dak-coloured moduble matter. The author found that a incurie of gene ten prespitated intacture of ferm chlorida bluich black, like nut-gells, while the timeture of black ten gave a green colour with row, puts as esteehed doss.

A modification of the permanganate process, which appears to possess some advantages for the examination of tea, has been described by P. Dvorkovitch (Ber, xxiv. 1945, Jour Chem Soc, lx 1302), who aims not only in estimating the tannin but also the proportion of fermentation-products formed in the process of fermentation to which black tea has been subjected. For this purpose he treats 10 grammes of finely-powdered tea with three successive quantities of 200 c.c. of boiling water, five minutes being allowed for each digestion. The residue is then boiled twice with 200 cc. of water, or until the last extract is almost, if not entirely, free from colour, when the decoction is diluted to 1 litre. Forty c.c. of this solution is then diluted to 500 cc with water. and treated with 25 cc of indigo-carmine solution and 25 c.c. of dilute sulphune acid (200 grammes of H.SO, per litre). The liquid is then titrated with a standard solution of potassium permanganate (containing approximately 2 6 grammes per litre), and of such strength that 130 cc are equivalent to 100 cc of decinormal oxalic acid (6 3 grammes crystallised acid per litre). The mode of adding the permanganate is important, and Dvorkovitch recommends that in the titration of the indigo-carmine 18 c.c. should be added at the rate of 2 to 3 drops per second, and the remainder at the rate of I drop per second; and that, in the titration of the tea solution mixed with indigo-carmine, 23 c.c. of the permanganate should first be run in, the addition continued at the late of 2 to 3 drops per second, and finally 1 drop per second added until the reaction is complete. If more than 38 c.c. be required, a small quantity of tea infusion should be used. To estimate the fermentation-products, 80 c.c. of the tea infusion is mixed with 20 cc. of baryta-water (containing 4 grammes of baryta

¹ Prepared by mixing 50 grammes of pure indigo carmine paste with water, adding 50 grammes of sulphure scid and 1 lite of water, filtering, and diluting fill 25 c.c. require 20 c.c of the strukture first past water.

per 100 cc.), the liquid filtered, and 50 cc. of the filtrate (representing 04 gramme of the tea) diluted with 500 cc. of water, mixed with 25 c.c. of dilute sulphuric acid and 25 of the indigocarmine solution, and titrated with permanganate. 18 cc should be run in first of all, then 2 or 3 drops per second added, and finally I drop per second till the end of the reaction. The volume of permanganate required, less that reduced by the indigo solution, represents that required for the oxidation of the fermentationproducts of 04 gramme of tea. According to Dvorkovitch, the joint weight of tannin and fermentation-products is obtained by multiplying the weight of exalic acid equivalent to the measure of permanganate required for their oxidation by 313, since 63 grammes of exalte acid correspond, according to Dverkevitch's experiments, to 31 3 grammes of tea-tannin (as compared to 62 3 of querestannic and !). Employing this process, he found from 8 84 to 10 55 per cent of tanmn, and from 0 90 to 1 88 of fermentationproducts, in teas of the first crop of 1890; and he concludes that the higher the proportion of caffeine to the total amount of tannin and fermentation-products, the more valuable is the tea.

The Lowenthal process distinguishes the tannic acid from the small quantity of gallic acid also present in tea, but as the astringent character of the infusion is due to both these substances, a method which will estimate the total amount of astringent matter, without distinction of its nature, is in some respects preferable to a process that gives merely the amount of tannin, while ignoring the gallic acid Such a process was devised by F. W. Fletcher and the author in 1874 (Chem News, xxix, 169, 189), and was based on the precipitation of the tea infusion by lead acetate, and the use of an ammoniscal solution of potassium ferricyanide to indicate the complete precipitation of the astringent matters. In practice, 5 grammes of neutral acetate of lead should be dissolved in distilled water, and diluted to 1 litre, and the solution filtered after standing. The indicator is made by dissolving 0 050 gramme of pure potassium ferricyanide in 50 cc, of water, and adding an equal bulk of strong ammonia solution. This reagent gives a deep red coloration with gallotannic acid, gallic acid, or an intusion of tea One drop of the solution will detect 0 001 milligramme of tannin, or 0 001 gramme dissolved in 100 cc of water. In carrying out the process, three separate quantities of 10 cc, each of the standard lead solution should be placed in beakers, and each quantity diluted to about 100 cc with boiling water. A decoction made from 2 grammes of powdered tea in 250 cc of water (the same as is used for determining the extract) is added from a burette, the first trial quantity receiving an addition of 12, the second 15, and the third 18 cc., or if green tea be under exmunation, 8, 10, and 12 cc may be preferably employed Portions (1 cc) of those trial quantities are passed through small filters, and the filtrates tested with ammoniacal ferretyamide solution

The approximate volume of tea decoction required in this easily frond, and after repeating the test nearly the nequate measure can be at once added. In this case about 1 c c of the liquid should be removed with a papetr, assend through a small filter, and diops of the filtrate allowed to fall on to apots of the indicating solution previously placed on a provedient slab. If no pank coloration is observed, another small addition of the tea decoction is made, a few drops of the hupd filtered and tested as before, and this process tepeated until a path colour is observed. The greatest delication is obtained when the dops of filtered solution are allowed to a obtained when the dops of filtered solution are allowed to proint of unchoos of the hundred of observing the rount of unchoos of the hundred.

The volume of tea solution it is necessary to add to 100 c.c of pure water, in order that a drop may give a pink reaction with the indicator, should be subtracted from the total amount run from the burette

The foregoing piocess is simple, and gives very concordant results, but the repeated filtrations required for the observation of the end-reaction are opt to be tedious. It is difficult to obtain pure tainin for estating the lead solution, and hence it is preferable to abundom the attempt and make pure lead sectate the startingpoint. The author found that 10 c.o. of the lead solution would prespitate 0010 gramme of the pricest galletaniar acid he could obtain. Hence, it all the weights and measures above mentioned be adhered to, the number of c of tea decoution required, divided into 125, will give the pencerings of Lumin and other precipitable untates in the sample. The proportion found in undired black tea by F W Betches and the author ranged from 85 to 115 per cent; with an average of 10 per cent. A sample of howas entechnic tested by the lead process gave a result corresponding to the presence of 119 per cent of taining (ac). (Son also page 491)

Another simple method for the deteninanton of tannun is that of J M E de x (Dunyl Folly Tour, coxxxx, 81), which consists in precipitating the boiling desoction of 2 grammes of tea with access of a 5 per cent solution of capure access the The precipitate is separated by filtation, washed, dred, and ignited. The resultant captine canle, GuO, can be moistened with turne acid, is-figured and weighed as such, or re-gratted with sulphur in a closed cracible, and thus converted into an equal weight of non-hygroscopic cuprous sulphide, Gu₂S. The weight obtained, multiplied by I 305, gives that of the tanum precepticated. The method is said to give

results correct to within 02 to 03 per cent; but any pectous bothes should be previously separated, if present in quantity, by precipitation with alcohol. By this method Edei found an average of about 10 per cent of tanini in twenty-five samples of black tan, and 12 to 12½ in green and yellow ten. S. Janke, by the same process, found from 6 9 to 91 per cent of tanini in black tan (origitates assumples), and 8 to 9 un green. Cupric acetate may be extemporised by mixing a solution of cupric sulphate with excess of sodium acetate.

C M Cames (page 491) obtained results by Eder's method closely agreeing with those yielded by the same samples with the lead process, and hence the proportion of callic acid in tea is

probably very magnificant.

In the case of caper and he teas, the astringency is often very high. owing to an admixture of extraneous fannin matters, but the evidence of the presence of such additions afforded by the determination of the tannin of tea is, of course, merely inferential Strong infusions of genuine tea, with the exception of some kinds from India, are generally quite clear, and do not become muddy on cooling Tea adulterated with catechy gives an infusion which quickly becomes turbid on cooling. More direct proof of the presence of catechu may be afforded by the following test devised by the author, which, however, should always be applied to the suspected tea side by side with a genuine specimen .-One gramme of the pure tea and an equal weight of the suspected sample are infused in 100 ce, of boiling water, and the strained liquid precapitated while boiling with a slight excess of neutral lead acetate Twenty cc of the filtrate from pure tea (which should be colourless) when treated with a few drops of silver nitrate solution (avoiding excess), and cautiously heated, gives only a very slight greyish cloud or precipitate of reduced silver, but the same tea containing 2 per cent of catechu (purposely added) gives a copious brownish precipitate, the liquid acquiring a distinctly vellowish tinge With a somewhat larger proportion of catechu, the filtrate from the lead precipitate gives a bright green colour on adding one drop of dilute ferrie chloride, while the solution of pure tea gives only a slight reddish colour due to the presence of acetate On allowing this liquid to stand, the adulterated tea gives a precipitate of a greyish or olive-green colour, the pure tea undergoing no change.

These tests, which depend on the properties of estechnic and, together with the excessive proportion of astringent matters (as shown by the lead process), render the detection of any considerable proportion of estechn tolerably certain, but a means of detecting mail additions is still a desideration.

Catechu is usually introduced in the forms of caper and lie tea, but appears to have been sometimes added in a separate state, to impart additional "roughness" or to cover the presence of exbounted leaves

Caper is a name applied to tea which has been made up into small glossy gammar masses by the aid of gam or starch. Some years ago the caper tea from the Centon district was invariable obtlierated with saidy and magnetic matter, and often with catchin or other extaneous astringents, together with foreign leaves. Notwithstanding the conviction of Wigner and some other authorities that genuine caper tea exists, the author believes to be invariable as features article.

Lie ice is the name given to a fraudulont mixture consenting and, weepings and dust of tea and other leaves, mixed with elay, and, uren one, &c., and made into irregular masses by means of gum or starch. When put into hot water, in the admiraction and falls to powder. The sodine test for starch may be applied aften and fulls to powder. The sodine test for starch may be applied aften and fulls to powder. The sodine test for starch may be applied aften and the property of the solid beginning with parmangenate. The sah of he tea is sometimes as high as 30 to 40 per cent.

The msoluble matter and extract of he and caper tea are very variable, but the former, exclusive of minoral matter, is usually considerably below the proportion yielded by genuine tea. The gum² in caper tea often amounts to 15 or 20 per cent, while the soluble sath so frem less than 2 per cent

The following figures show the results to be expected from the analysis of factations tea .--

	Α.	В	О
Observer,	J, Bell.	J M Ader	A B TIII
Description,	"Mahloo mixture "	Black tes	Green tea.
Extract,		22 40	87-00
Tannin, .		19-77	Catecht
Total Ash,	997	(Catechu detected) \$ 07	detected, 12 10
Magnetic and sandy matter,	4.81		6.00
Soluble Ash,	1.54	1 12	1.29
Alkalinity, at E ₂ O, .	0:17		0 18

At the present time (August 1892), Canton espers are frequently loaded with from 3 to 5 per cent of sand, &c., but they raisely appear in the home market, being stopped by the Customs, or purposely imported for future exportation (M. J. Shoridan).

The gam is determined by concentrating the squeous decoction of the tea

The following analyses of samples of apurious tea, recoved from the U.S. Consults at Canion and Nagsank, are by J P Battershall (Food Adulteration, page 28). No. 1 consisted of partially exhausted and refired leaves known as "olang seep" (clear water), as me appearently referring to the character of the unicon. No. 2 was a samter of "betoe" made from wampan leaves No 3 was a mixture of 10 per cent of gient eave with 90 per cent of lie-tea, sometimes sold as "Impaini" or "Gunpowler" teast made up into hittle shot-like pellets by means of "Congou paste" (Cooled no).

					No 1	No. 2	No 8	No 4
Insoluble leaf,		-	-	•	70 00	70 55	67 00	60 10
Extract (complete),					778	14 00	1276	22 10
Gum,					10 67	7 30	11 00	11 40
Tanuin, .					\$ 18	8 01	14 50	15 64
Caffeine,					0 58	none	016	0 12
Ash -Total,					8 62	890	7 95	12 58
Soluble in water	, .				0.64	1 88	8.00	8 84
Insoluble in acid	١, .				8 92	8 18	188	8-60

Logswood is mentioned by Eder as an adulterant of tea To the test the steeps the tea in cold water. If logwood he present, the resultant solution is changed to a bright green on adding a hittle sulphunc acid, and to blackish-blue by a solution of neutral chromate of potassium

Facings and colouring materials were formerly almost invariably to the object being to impart a fine demanded by custom but not naturally possessed by the leaf Colouring matters have been extensively employed for transforming black tea of low outlity into superior grean.

In the case of Roberts v. Egeiton, heard before the Court of

almost to an extract, treating the residue with strong spurt, and filtering and washing with epurt. The preceptate is sunsed off the filter with hot water, end the colution evaporated to dryness at 100° The residue is weighed, ignited, and the ask weighed. The loss is regarded as "grup," but is lable to be in excess of the truth from the presence of albuminous matters

¹ It is a fact well known to the trade that for many years a certain firm of tea merchants used some method of removing the facing after the arrival of the tea in the country. Queen's Bench in 1874, Mr Justice Blackburn decided that the facing of green tea with gypsum and prussian blue was an adultation, because natural green tea could be obtuned without such means.¹

If a faced tea he examined under the microscope as an opaque object, the nature of the facing materials may often be recognised. On beating a faced tea with warm water, the colouring matters become detached, and the small portions using to the surface may be floated on to a giass shids and at once examined under a microscope, while the bulk of the facing is obtained as a sediment when the stanned luqud is allowed to stand ²

For sign leaves in tea are legitimately present in small proportion (1 to 3 per cent) to impart bouquels fut larger admixtures can simply be regarded as due to adulteration Slos, older, and willow leaves have been (formerly) met with in England as adultionates of teat. Among the recontly-found leaves added abroad, and stopped by the Customs, are those of Olito authus inconspicius, Camellia seagangus, Euro Chimensa, and also see In the recognition of foreign

 1 The tass consumed by the Chinese and Japanese then selves are not faced. According to Y Kozai the maximum proportion of faring in the green tea of Japan is about 0.4 per cont

I This deposit often has a distinctly greenals colors from the presence of pressure from pressure from the deposit of the pressure from the first state of the deposit of t

⁸ As a rule, the odourferons leaves are not allowed to remain in the tea, but having imparted their characteristic flagrance to the tea are removed previously to packing.

⁴ From the result of a parlamentary unquiry hold in 1885, it apposed that payends of four multino pounds of actions to swe so animally prepared in this country from sloe leaves, and used to adulterate Okina tos. Up tall within a key seas of that date this hilten practice was centred on accessly, but subsequently a patent was obtained for the preparation of thirth leaves as a substitute for test, and an extensive manufactory was established for this purpose. The industry was ultimately suppressed, and a large quantity of the product horizon!

b In 1888 Wends and Wioregerski found in the tess sold in Warsaw various foreign leaves, which they identified by their austomical characters. leaves in tea, chemistry cannot be expected to play a very active part, though it sometimes affords very useful indications. Thus A. Wynter Blyth has pointed out (Analyst, ii. 39) that a crystalline sublimate (which he believes to be thorne) is obtainable from a single leaf of tea. For this purpose he boils the leaf for a minute in a watch-class with a very little water, adds an could bulk of calcined magnesia, and ovaporates the mixture ispudly to a large diop. which is transferred to a microscopic covering glass and evaporated nearly to dryness on a heated from plate. It is then covered by a ring of glass, and when the moisture is nearly driven off a second slip of glass is added as a cover At a somewhat higher temperature theino volatilises, and on examining the deposit on the covering under the microscope may be recognised by its characteristic appearance Other leaves than tea may give a crystellino sublimate, but if no sublimate is obtained the leaf cannot be a product of the tea-plant.

A W Blyth has also proposed to utilise the constant presence of manganese in tacheaves as means of recognising them If a single tea-leaf be ignited in platinum, and the sah taken up in a best of solution emborate contained in a boop of platinum ware, on remeding the flux after a munite addition of into the green colour of the solution manganate will be distinctly necognisable Or a munite quantity of intre and earlomate of sodium can be at one added to the sah on the platinum fail, when on fruing the mixture a distinct green colour will be obtained if mananese to measure

The author has found mangenose in the leaves of Canellata Thea (case), Canellata Approince, Canellata assengine, Coffee As abuse, beech, blackberry, and symmore. Mangeness was absent from the leaves of the hawthorn, ash, is spherry, cherry, plum, and rose, and only faint times were detected in the leaves of the Ree Paragrapheness, elin, butch, lune, slos, elder, willow-hesb, and willow

For the detection and identification of foreign leaves in tea, the botanical and minoscopical characters are best fitted. Some of the sample to be examined should be put into hot water, and Among the larves recognised was those of Biplohums angust/slum, of Bicube will or \(^1\) to \(^1\) which-formed the great part of the "tax" sold in outsia localities. They also found the leaves of Epidohum harmston (great wildown-berry), Francisco acceleror (sub), Sambouss supa (clobe), Pagarar sone all conventions), Francisco acceleror (sub), Sambouss supa (clobe), Francisco acceleror (sub), Sambouss supa (clobe), Tomos consine (degree), and zides approxim (Date Carriard). The inlusion of wildow-has be derived than that of its, and gives a preceptate of muclage on transment with alcohol. An article known in Blassa is "Karpat Las" also contains an admixture of the larves of Epidohums angustificium. Two samples examined by Ji-Nittinisky in 1885 yielded 78 or and 10.45 per cent of ash, are rest of ash, are 1885 yielded 78 or and 10.45 per cent of ash, are rest of ash, are sent of ash are sent of ash are sent of ash, are sent of ash are sent of

sentative genuine tess yielding from 5.60 to 6.87 per cent

when the leaves have unfolded they are spread out on a class plate and held up to the light, when, with the aid of a lens. the venation, serration, &c , can be readily observed A valuable aid to the examination consists in treating the leaves with a solution of sodium hypobromite, or, as suggested by A. Wynter Blyth, a. strongly alkaline solution of potassium permanganate. In using the reagent, the leaf should be enclosed between two microscopic cover-glasses, a weight being placed on the upper one to keep it in position. On heating the leaf with the reagent, action at once commences, the colouring matter being first attacked and subsequently the cell-membranes. When the action is sufficiently advanced, the leaf is removed, washed, and immersed in hydrochloric acid, which leaves the leaf as a translucent white membrane in which the details of structure can be readily observed J Bell removes the skin of the leaf by immersing it in "water containing a few drops of nitric acid," and gradually heating to the boilingpoint, when the skin rises in blisters, and may be readily removed. by a camel's-hair brush

The primary venation of the tea-leaf consists of a series of well-defined loops, which are not met with in most leaves hickly to be used as adulterants. The seriations are not more saw-teath on the margin of the leaf, but actual hooke. The seriations stops short adruptly at some distance from the bess. The Assum tea-leaf is semestimes baserate. At the apex of the tea-leaf there is a distinct note, mested of a point. The epidermus of the under-surface is seen under the microscope to consist of distinct amounts cells, with numerous oval stomats, and a few, long numerous. If the under-surface of the tea-leaf be examined under the microscope after separation of the cutific, the peculiar and characteristic space between the twincells of the stomata may be readily precisived.

T Taylor has pointed out the presence of "stone cells" in the leaves of tea and Canalità Apponica, and confirms the observations of Blyth as to the absence of these formations in the leaves of the willow, aloc, beach, ash, black-currant, nephenry, and Ex-Paraguagensis Taylor prepares the leaves for examination by boiling them in a strong solution of caustic points or sola.

¹ The serrations are very strongly marked on mature leaves, but are indistinct or almost wantang in the delicate leaf-buds which constitute "flowmy pickoe."

² Tea-hairs are comeal, pomted, slightly bent towards the base. They have very thick walls, and the central duct usually contains granular matter. Numerous hairs are observable on young tea-leaves, but on old leaves they are sometimes wholly wanting.

In the leaf of the blackthorn or slos (Pranus communic or P. spinsso)! the scratures are direct incisons, numerous, often irregular, and extending to the base. There are no spines. The cells of the spidernus are not amous, and are much smaller than those of tea, especially on the under surface. The cells on the upper surface are strated. The stomata of the sloc-leaf as smaller and less numerous than those of tea. The hairs are shorter and coarser than those of the tea-leaf, are marked in a peculiar manner, and have a club-baned calarterment at the base.

The leaf of the elder (Sambucus myra) is more pointed than that of the tea-plant, and the lobes are unequal at the base. The serrations are direct incisions. The midriff has hairs on it, and on the leaf itself there are two distinct kinds of hairs—one, a short,

spinous hair, and the other jointed and club-like

In the leaf of the willow (Scilic alba) the scratnons much resemble those of tas, but the cells of both the upper and under epiderms are much smaller than in tes, and the walls are not smuons. The hans, which are very abundant on both actes of the leaf, are long, uncellular, and smuous The elongated form of the willow-leaf and the character of the vension also distinguish it from tes

The appearance of the leaf of the hawthorn (Crategues monegque and C. expacanital) is well known. The cells of the epidermis are mostly quadrilateral, with very sinuous outlines, especially on the mider surface. The stomata are oval or nearly round, large, and numerous.

The leaves of the beech (Fagus sylvatica) are ovate, obscurely dentate, with parallel venations lunning right to the edge

The leaves of Chicronthus suconspicuse are long, o'al, sorrated, wrinkled, with venations running nearly to the edge, and there by their intersection forming little knots which give the margin of the leaf a very rough feeling. The cells of the epidermis are very larce, and the stomats oval and rather numerous.

The leaves of Camellia escangua are oval, only obscurely serrate if at all, and of a tough leathery texture. The lateral venus reinconspicuous. Both the upper and lower epidermis show a peculiar dotted or retaculated structure, and the lower is studded with numerous small obloing stomate.

. The leaves of Lithospermum officinals (the common gromwell) have been extensively used in Bohemia for adulterating tea. They

A specimen of slos leaves gathered early in September gave, after drying, the following results (in the author's laboratory)—Mosture, 6:40 per cent, insoluble matter (oh whole leaves), 55 90, tannin (by gelatin), 18 00, gum, &n. 8 90, total sels, 8:74, and ash soluble in water, 4:70 per cent,

are lanceolate, with a harry under-surface, are destitute of alkalord and essential oil, contain about 9 per cent, of fat and 8 of tannin, and leave about 20 per cent of ash on ignition (Jour. Chem Sec, xl 131).

The general appearance and venation of tea, and leaves which have been, or may possibly be, employed for its adultication, as shown by two plates at the end of the volume (page 572). The illustrations are life-size reproductions, by the collection process, of photographs of leaves, taken by J. T. Stevenson in the author's laboratory

A Wynter Dlyth has pointed out the churacterists on power ance of the "skeleton-ash" left on iguiting leves from different sources. The lenf to be examined as piaced between two cucles of microsopic covergings, the upper one woughted with a alver con, and the whole ignited cuttously in a flat platinum distior on platinum fol. Before the earbon is completely consideration of the continued, and the skeleton-ash examined under the microscope.

Mate. Paraguay Tea.

Mate1 or Yerba consists of the prepared twigs and leaves of liew Paraguayensis, or Brazilian holly 2

Bysson found in e.a.f.-gu a.e. u, the commonest kind of mate, consisting of the large and old lowers with twings and fingements of wood —Caffeine, 185 per cent, a substance resembling bit-lime, fatty and colouring matters, 387; complict glucoside, 288, resm., 683, mineral matter, 392, and an undetermined proportion of males and matter, 392, and an undetermined proposition of males and matter, 392, and an undetermined proposition of males and matter, 392, and an undetermined proposition of males and matter, 392, and an undetermined proposition of males and matter and

Some fresh leaves of Res Paragangassis, grown in Cambridge Betained Gardens, were found in the author's laboratory to contain 681 per cent of water An analysis of the same leaves after drying at 100° C. showed —Insoluble matter, 57°94 (= hole-vater extract, 42°04), tamin by "Th\$_1 16°15, tamin by Cu\$_1 16°06; ceffous, 118, total ash, 614, soluble ash, 3°56, alkelinity of soluble ash (as K,0) 0.712 per cent.

A. W. Hofmann found in mate 0 3 per cent of caffeine and a variety of tannin identical in every respect with that present in toa.

¹ The word mate is not accented, as sometimes written, but it should be pronounced as two syllables.

⁹ Yanonta alliad speness are recognised, but Hee Reagnagements appears to the oho only one cultivated. It has been grown in Spain, Fortugal, and Cape Colony, in addition to its native habitat. At the present time it is used by about 12,000,000 of goods, the annual consumption in the Argentine Republication being twenty-area million pounds.

P N Arta found the tannum of mate to be analogous to but not identical with that of coffee On dry distillation he found it to yield resorting as well as sateshol Caffestimms such he regards as dioxy-parasimansylte and, and matelanine and as belonging to the group of exprincylinguous and Southerram and Delondre state that mate contains the same essential constituents as the coffice-leaf, and in greater proportion than the coffice-such size of the work of the subject (Phan m Jour, [3], xiv 121), including some elaborate proximate analyses of mate

The aromatic principle of mate has not been isolated, but by dry distillation a volatile oil of phenolic character is obtained

The ash of mate resembles that of tea in containing a notable proportion of manganese

The leaves of the Yopon (Rec cassue), a shub or small tree growing on the coast of Vingma and Carolina, have been used as a beveauge 1 F P Von able (Rhem News, lu 172) found in an an-driet sample —Mosters, 1819, water extact, 2655, tannin, 759, caffeine, 027, and ash, 575 per cent The sah contained manacases.

Coffee.2

Commercial coffee consists of the seeds of Oglica Aradaca and allied species belonging to the order Cinchonaces 8 The coffeeties is a shrub-like plant cultivated in various tropical countries. The best coffee that reaches England comes from India, Java, and Ceylon. A little "Mocha" coffee comes from Araba, but the

Although the leaves of ta, coffee, and Baznhan holly are almost the only one known to centum onfluen, a beweage sy nepson from the leaves of many other plants in various pasts of the world. Thus, Culhe white, a sketness related to the symulate, as extrawed venithread in the interior of Anaba, and the leaves, known as K h a, Coffa or Anaban taa, are used both as a between and for obsening Faburn, or oracle tas, is made from the leaves of Angreeous fragious, growing in the Mauntius, and some years mace was unloaded into Plans as a regular article of comment. The Ar a ho, a midstitute for too which has been sold in Para, counts of the small leaves of Piso computer for the oracle of Piso computer. The Pison growing on the shopes of the Adaps Mountains. Bas form or Trebison do ta a semide from the leaves of Viscansius arotaclyphylog, a plant clearly allulat to the canability Cap te can and Bast har discribed in the factories on page 503. Karpen taa sidescribed on page 523. "Nonthi, Left, off. German, or Knode."

⁸ Three species of Coffee, distinct from each other, are now grown —1. The Arabien or Mocha coffee-plant has short upught bianches, with a brittle leaf and seeds usually single in the berries. 2. The Jamiesz coffee-plant bears longer and more pivible bianches than the Alabian, has a toughel leaf, and the seeds are almost always double in the beries. 3. The East Indian or Description.

greater part from India Biazil at the present time furnishes about one-half of the world's supply of coffee 1

G o m m a 111 c (Monst. Scient, [18], vr. 779) gives the following the chemical composition of undressed Mysore orefite:—Moisture (from 24 samples), 63 to 157 per cent, fathy matters, 1298; glucose, 260, legrum-cassin, 152, albumin, 104, caffeine, 042 to 131, and sab, 388 per cent

O Levesie (Arch Pharm., [5], iv 294; Jour Chem. Soc., xxxi 752) obtained the following range of figures by the analysis of seven typical samples of raw coffee.—

Cafforne,					0.64 to 1 53 per cent.
Gumniy matt	er,				20.6 ,, 27.4 ,,
Fat, .					14.76 ,, 21.79 ,,
Tanns and co	ffetanni	c acids,			19.5 ,, 28 1 ,,
Collulose,					29 9 ,, 88.4 ,,
Ash, .			٠		8.8 ,, 4.9 ,,

J Bell (Analysis and Adulteration of Foods, i. 43) gives the following analyses of typical samples of raw and roasted coffee .—

Bengal plant has smaller leaves than the Jamanca coffco, and very small bernss. The Liberan coffee-plant (Coffee Liberan) appears to be a distinct species, which is little subject to discase, and has been successfully introduced into the East Indias.

The offee fruit usually, but not always (see above), contains two twin seeds, which touch each other on the flattened surface. These are contained in a pulp which is removed by water and a process of fermentation; and the membranous personary (seebnically termed "parohiment") which incloses sach seed is is more 4by 10 ilbra and wranowing.

The parchment from coffee-berries is imported to England in considerable quantities, and, when reasted, is said to form an ingredient of this beverage sold in cheap coffee shops

An analysis of unroasted "perchment," made in the author's laboratory by O. M. Caines, showed it to contain .—Water, 9 43; essential oil, 0 665; calfinis, 0.27, hot-water errors, 1.61, total ssh, 10 41, and soluble ash, 0.19 per cent. A somewhat coffee-like a come was doveloped by reasting

It is stated that the Arabs in the neighbourhood of Jedda discard the kernel of the coffee berries and make an infusion of the hueks (*Pharm. Jour.*, [3], xvi. 669.

In Australia, an infinison of slightly reasted coffee-leaves a dutuk in the same manner sets. Its tasts asquests at once that of tea and tobacco. The loaves, when burnt or roasted, exhals a powerful edour of tobacco, and the small of the condimented vapours strongly suggests that of tobacco-junes. O He'hr or, who has snalysed the leaves (Analys, xv. 84), found only 0.29 per conit of coffeen.

	Mochi	Coffee	East Indian Coffee		
	Raw .	Ronsted	Raw	Roasted	
Melature,	8 98	0.68	9 84	1 18	
Claffeine,	1 09	-82	1 11	1 05	
Saccharino matter,	9 55	48	8 90 `	41	
Caffero ands,	8 48	474	9 58	4 52	
Alcoholic extract, containing nitrogenous and colouring matter,	6 90	14 14	4 81	12 67	
Fat and oil,	12 60	13 59	11.81	13 41	
Legumin and albumin,	9 87	11 28	11.28	13 13	
Dextrin,	87	1 24	84	1 88	
Cellulose and insoluble colouring }	87 95	48 02	88 60	47 42	
Ash,	8 74	4 58	8 98	4 88	
	100 00	100 00	100 00	100 00	

Bell beheres the sugar of coffes to be a peculiar species, possibly aliaed to melentose G L Spenos, on the other hand, has definitely proved that the earbohydrates of coffee conast very largely of sucross, which he has notated on considerable quantities. There is hkowise present a body which yields galactose on hydrolysa, as also a peutoser-picking gum

CATPMANTIO ACID, \hat{C}_{0} Ha $_{10}\hat{C}_{0}$ called by Payen ch lorge an is at all exists in offse-betres in the proportion of 3 to 5 per cent, probably as a calcium or magnesium and caffeine. It is prepared by diluting an alcoholic infusion of offse with water, filtering from precipitated fatty matter, and precipitating the boiling filtest with lead acetacle? On decomposing the washed precipitate with sulphuretted hydrogen free caffetamic acid as obtained. It forms a yellowish-white powder, or groups of colouriess manufallated

1-W. H. Krug determines caffetanue and as a lead sait. He treats 2 grammes of coffee with 10 a. of wets, and dispests for 38 hours, then add 25 c. of treatfield spurit, and dispests and the sense of treatfield spurit, and the distribute heatest to the boultage rout. A bouling concentrated solution of lead sociate is added, which throws down a prespiration of Po(QuH,Qo). When this has become flocusing the lifetime of the distribute and the state of the distribute of the dis

erystals. It is very soluble in water, less soluble in alcohol, and only very sparingly in ether Coffstanine and has an estimagent taste, and the solution strongly reddens litanus. It gives a dark green colonation with forme chlorate, and precipitates the sulphates of quinne and encohome, but not gelatin, ferrous salls or intanemetic. It rediness silver mitrate on heating, forming a metallic mirror. The salits turn green in the air.

On dissolving calificatine acid in caustic alkali or aumonia, and exposing the solution to the air, the highd acquires a bibush-green colour owing to the formation of the oxidation-product, viridic acid, which is an anonphous brown substance, very soluble in water to form a solution which is turned green by alkalies. It gives a blumb-green precipitate with haryta-water, and a blue with lead acetate. Viridic acid dissolves in oncentated subjunct and to form a crimson solution, which on dilution with water gives a foccellent blue precipitate.

On prolonged boiling with caustic alkalies, caffusianne and is split up mile a sugar and caffore a crid $Q_a F Q_{c,0}$ which crystallises from the neutralised liquid and has the constitution of a dhydroxy-emmanue and Dy fusion with caustic potais), caffetamic acid yields protocatechuic and acetic acids. Hosted alone it gives catechol

ROASTING OF COFFEE During the process of roasting, the aroma of coffee is developed and the toughness of the beam destroyed, so that subsequent grinding is facilitated. If the roasting be insufficient, the namess is not destroyed and the flavour

not fully developed, while if over-roasted, the product has a nauscous empyreumatic flavour

When resisted to a yellowish-brown, coffee losses, according to Cade is, about 12 per cent of its weight, and in this state is difficult to guind When reasted to a chestrut-brown it losse 18 per cent, and when it becomes entirely black, though not all earbouned, it has lost 23 per cent. In practice, the less of weight in reasting coffee is between 12 and 20 per cent (of which about 8 per cent represents water removable at 100° C), and if the latter figure is reached, the product is injured. According 10 W at so in W 11, the usual yield of reasted coffee is about 98 flow from 1 cut of raw berries. This corresponds to a loss of 12 5 per cent

K on 1g found that on resating coffee-herries to a light blown the total less of weight was 17 77 per cent, of which 866 was water and 911 per cent organic matter. The original coffee contained 11 19 per cent of moisture, and after rossting, still retained 310 per cent. Eliminating this extraorective water from the results.

the percentage composition of the raw and roasted coffee was as follows:—

				1	Raw	Roasted
Caffeine,				_	1.88 per cent	1 42 per cent
Fat,)	14 91 ,,	16 14 ,,
Allominous ma	ters,			1	11 43 ,,	12 31 ,,
Sugar, .					3 68 ,,	1 85 ,,
Undefined non-r	itrogenou	nat	ters,		84 55 **	39 84 ,,
Cellulose,				- 1	81 24 ,,	25 07 ,,
Ash,			٠	-	8-92 ,,	8 87 ,,
					101 04 (f) per cent.	100 00 per cent
Total matters soluble in water,					30 98 per cont	28.86 per cent.

According to Paul and Cownloy (Pharm. Jour., [3], xvn. 655, 821) there is no appreciable loss by volabilisation of caffeine during the reasting of coffee, unless the process is carried to excess But Paul admits that the water condensed in the place leading from the roasting often contains some caffeine, which he considers has been probably carried over mechanically (Pharm Jour., [3], xvn. 231). Wat son Will (bdf., page 684) states that he has never failed to find caffeine in the sublimate obtained in coffeerosating.

The chemistry of the roasting of coffee has been studied by O. Bern heimer (Monatsh. Chem., 1. 456, Jone Chem. Soc., xln. 230), who roasted coffee till it had lost about 25 per cent of its weight? The uncondensible vapours consisted chiefly of carbon

¹ Paul points out that the caffeine exists in coffee in the form of caffetannato, which compound will not suffer decomposition at the ordinary temperature of reasting. Considering the great facility with which salts of caffeine undergo decomposition, this statement seems to require confirmation.

* Prily kologrammes of onfice yacidad is hitzer of aspasses sitellate and 68 grammas of solal matter floating theoron. On a spitzing the with either, sitty acada, quanol and caffool wave extracted, while caffones, acotte acad, mothyl mane and tumethylamine remanule on the capteons haped On evaporating the etheresi solution, and fractionally relating the restead dark, code-smalling of a few days of a section-this liquid passed over, followed by a little socitic of the contraction o

dioxide, and by passing them through dilute hydrochloric acid a resinous substance having the appearance of pyrrol-red was denosited. Among the solid and hould bodies volatilised. Bernheimer found .- Palmitic and other solid fatty acids, 0 48 per cent , caffeine, 0.28 per cent ; caffeol, 0.05 per cent , besides water and acetic acid. Quinol, pyrrol, acetone, methylamine, and trimethylamine also occurred as secondary products

CAPPEOL, C.H.,O., is an only liquid smelling very strongly of coffee, and no doubt is the substance to which the aroma of roasted coffee is due. It may be obtained by distilling roasted and powdered coffee with water, shaking the distillate with other,

and evaporating

Caffeol boils at 196°, and is not solidified by a freezing mixture. It is not sensibly soluble in cold water, to which, however, it unparts its characteristic odour. It is slightly soluble in hot water. very slightly in aqueous potash, and with great facility in alcohol and other The alcoholic solution gives with ferric chloride a red coloration, said not to be destroyed on adding sodium carbonate. By fusion with caustic potash, caffeol yields salicylic acid, and, according to Botsch (Monatsh Chem, 11 621, Jour Chem. Soc., xlti, 174), is isomeric with methyl-salicyl alcohol, the two compounds having the following constitution :--

C.H.(O CH.) CH.OH Methyl salicyl alcohol

CaH,(OH),CH,,OCH,

Paul and Cownley (Pharm Jour, [3], xvn 822) found that on infusing coffee in six times its weight of boiling water, about 88 per cent of the caffeine passed into solution. Three fluid ounces of such an infusion contained 2 36 grains of caffeine As the medicinal dose of caffeine is from I to 5 grains, a cup of coffee may be expected to have a marked effect as a stimulant

The dietetie value of coffee is possibly dependent as much upon the presence of caffeol as on that of caffeine. According to M. Fargas, the effect of caffeol on the heart's action is the opposite to that of cafferne, and increases the strength and randity of the pulsations

According to Couty, Guimaraes, and Niobey (Compt. Rend , xoix 85) coffee diminishes the activity of the simple combustions which produce carbon dioxide, but increases the forma-

which was separated from the aqueous solution of soap and washed with water containing a little caustic alkah This dissolved out quincl, which was isolated by acidulating the washings and extracting with other. The viscid oil, consisting of impure caffeol, was dried by calcium chloride and fractionally distilled, when the greater part passed over between 195° and 197°.

ton and excreton of urea, and the assumilation of meat and other untrogenous foods It is a complex alment which renders the organism capable of consuming and destroying larger quantities of untrogenous substances, and hence may be regarded as an induct source of available energy

Commercial coffee is subject to a variety of sophistications, both in the form of herry and after grinding. So far as the United Kingdom is concerned, the majority of the frauds formerly practised are obsolete, or nearly so, but certain illust practices subsist

COFFEE-BERRIES vary considerably in size and character according to their origin. The following table shows the number of seeds required to fill a 50 c.c. measure (Thorpe's Dict. Applied Ohem., in 578).—

```
Fine blown Java,
                          187
                                Good ordinary Java. .
Fine Mysore. .
                          198
                                Fine Cevion plantation.
Fine Neilgherry.
                        . 208
                                Good average Rio. .
                                                            286
Costa Raca.
                           208
                                Medium Cevion planta-
Good ordinary Guatemala,
                           207
                                   tron
Good La Guayro,
                          210
                                Manilla, .
                                                            248
Good average Santas.
                           218
                                Ordinary Mooha.
                                                            270
Fine long-berry Mocha.
                       . 217 West African. .
                                                            313
```

' (Bull Soc Ottom, xivus 501), raw coffee by sea-water is sometimes weahed, de-counsed with nun-wave, again weahed, dred xandly, and coloured either by slight roasting or by dyeing with accoranges. By such manipulations, green Santae coffees are sault to be increased 25 per cent in value, and made to pass for Java growths E Waller states that South American coffees are often exposed to a high moist heat, which changes their colour from green to brown, in ministance of Java coffee. He found coffee-berres coloured with Scheele's green, yellow other, chrome-yellow, burnt number,

¹ We set Indian coffee-bernes are negulat un ass, pale yellowsh, firm and heavy, with a me arona, and they lose compassively little on reserved. Heavy, the me arona, and they lose compassively little on reserved as a "low" or "O're midding." ¹ as a nee oo folke between as unasile, plassed elongated, light, and deficient in aroms and essential oil. When new, Java coffee is pale yellow, and of less with than when old on brown. The description of the coloring is pale yellow, and of less with than when old on brown. The description of the coloring pale well as age. It has been artificially solouted to Oyi in profuses all descriptions of coffee, but the ordinary plantation coffees are even-coloured, slightly cance-shaped, strong in aroms and flavour, heavy, and more assempthie of adultmentant than the other kinds Gentlem Mocha coffee is small and dark yallow in colour, and considered of the hapster quality.

venetaan red, &c When possible, such facings should be detached by agitating the bernes with cold water and examining the sediment Organic colouring matters can be desected by soaking the bernes in alcohol, which is not coloured by genuine coffee On evoporating the alcohole solution to dryness, and taking up the residue in water, a solution will be obtained which will give the characteristic reactions of the coal-kut dyes.

The specific grantly of twenty-four samples of genume rawcoffee-hernes was found by Padd to range from 1368 to 1041, while the density of the same samples, after rosstarg in the either manner, varied from 0 635 to 0500. Baw coffse which is lighter than water may be suspected of having been damaged by sea-water or other means, and subsequently washed and improved in colour live artist possible.

The specific gravity of coffee-berries is ascertained by Padé by a special apparatis described in his paper. In the case of unreaded coffee, the gravity can be readily observed by immensing a few of the berries in saturated brain, which is then dulated with water till the coffee remains suspended in the liquid, the specific gravity of which is then taken. With nossels coffee, the brine must be replaced by the very lightest gasolene, the density of which can be increased if necessary by the gradual addition of ordinary kenesine. Another plan of ascertaming the specific gravity of coffee-berries is to introduce as many as possible into a tared 50 cc. flask or other vessel of known capacity. The weight is then ascertained, and the flask filled to the mark with mercury. The weight is again observed, when the increase will be the weight of mercury required to fill the intertaces between the berries.

Weight of berries in grammes × 12 59

(Measure of vessol in o c × 13 59) - weight of interstitual = sp. gr of berries, mercuny
mercuny

According to J Konig (Zestsch angew Chem, 1888, page 880) coffee is often roasted with an addition of glucose-syrup, which makes the decoction look stronger, and causes the beries to hold an additional 7 per cent of water. L. Padé states that roasted

1 Coffee so treated yealsh from 6 to 8 per cent of soluble matter on against with cold water, while offer onsated without sugar yaids from 6 to 5 per cent only. In the former case, Febling's solution indicates from 1 to 1½ per cent only. In the former case, Febling's solution indicates from 1 to 1½ per cent of reducing signar, against 0 2 to 0.5 in genuine coffee. Situater and Ratinari detect glucose by releasily against 20 grammes of the coffee heaves with 800 o. of water for first munter. The lunqui suffered indicate to 1000 oc and 80 oc of the filtested hugand evaporated to dryress at 100°. The dry readous we weepled, ignaried, and the said adouteds. Pur crosted

coffee-beans can be made to take up nearly 20 per cent of water by steaming them and coating them with glycenn, palm-oil, or vaseline to prevent-oraporation. The specific gravity of the berries is theneby raised to 0 650-0 770, and hence is sensibly above 0 635, which is the maximum forms for senium coasted berries.

Van Hamel Roos (Revise Intern des Falesfications, May 15, 1891) has called attention to an ingenious method of sophisticating offee-berries. A sample examined by him had the nucroscopic sizucture of genume coffee, but showed an almost entire absence of fat globules, and gave an other-extract of less than 1 per cent. (instead of 12 to 14). Roos suggests that the berries had been used for preparing coffee-extract, and then re-ros-sted with addition of a httle sugge.

As a coating for coffee, T. W Moore has patented (Eng Pat., 5038, 1889) a mixture of milk or condensed milk, ground or powdered glue, "liquid glyeerin," and refined lard, with the addition in some cases of bicarbonate of soda, fine sait, and vinegar!

Imitation coffee-bervies were formerly manufactured of fine-clay. These were nucled with genume berries and roasted with them, when they absorbed some of the colouring matter and oil, and so remained a close imitation. On breaking such spurnous berries the colour would be seen to be principally on the exterior. The determination of the total sah and sines would at once lead to the detection of such a fraud

In 1850, Messrs Duckworth of Liverpool took out a patent for moulding chicory into the form of coffee-bernes, and quite recently several kinds of factitious coffee-beans have been described.

A factory for the manufacture of unitation coffee-berries on the scale of 40 to 50 kilogrammes daily was necently sensed at Lille by the French Government II appeared in ovidence that the composition of the product was —Checory, 15 kilogrammes, flour, 35 kilogrammes, ferrous sulphate, & kilogrammes.

Factitious coffee-beans recently seized in Roumania consisted of coffee-grounds, chicory, and peas.

In Åmerica there are several firms which extensively manufacture untation coffee-beams and "offee-pellets". These preparations usually comest of wheat-flour, chicory, bran, and occasionally coffee. Samples punchased and examined by the cliemists of the U.S. Department of Agriculture gave the following results:—

coffee shows from 0.44 to 0.72 per cent of soluble organic matter, and gives a solution only faintly coloured, but if iosated with sugar or glucose the organic extract ranges from 1.81 to 8.18 per cent, and the hquid is more or less strongly coloured

Appearance	Specific Gravity	Composition,
Rousted beens, Rousted beens, Rousted beans, Rousted beans, Rousted pollets, Rousted pollets, Rousted pollets, Raw beans, Raw beans, Light coloured beans, Rousted beans, Rousted beans, Rousted beans, Rousted beans, Rousted beans, Rousted parantee, Rousted granules, Rousted granules,	1 196 1 198 1 111 1 119 1 183 1 198 1 211 2 174 1 184 1 118	Wheat four voltes and chlorry "Knutk Kriffee" Wheat four, coffee, and chlorry "Knutk Kriffee" Wheat four, but, coffee, and chlorry Wheat four, knut, nod probably rye. Wheat four and coffee Wheat four and probably awdust, Wheat four and probably awdust, Wheat four and probably awdust, Wheat four the colores East and probably awdust, The color and probably awdust, Wheat four the colores East and probables Fee built and beat

A W. Relnatrom (Emp Pat, 14,970, 1889) has described a substitute for coffee prepared by boiling down whey or milk in a vacuum to a pasty consistency, forming the product into cakes, drying it below 100°, cutting it into pieces the size of coffee-beans, and reasting

L Jaunnes, in 1891, examined a factitious coffee consisting of acorns and cereals

An imutation coffee examined by J. Konig (Zeitzel, angen. Chem., 1888, page 580) doedly resounded real coffee in appearunce, but all the bernes were precasely the same shape. Under the immovacopie, wheat-statch was detected, and Konig concluded that he article consisted of rosasted wheat dough of low quality E. Fricke (Zeitzel angen. Chem., 1889, page 310) has described a factitious coffee containing caffeine, and apprentily made from imprince-seeds (compare page 544). K. Portel e (Chem. Centrallul, 1890, page 135) has described factitious Goffee-beans sold under the nume of "Kunst Kaffee," The following were the compositions of the samples referred to above —

	Portèle	Honey .	Fricks.
Moisture, Proteids, Ret, Ret, Ret, Staroh, sugar, guso, dec Celliniceo, Caffeine, Ash,	1 46 per cent 13 93 " 8 96 " 64 01 " 15 83 " 0 07 " 3 13 " 101 63 per cent	514 per cent 1078 32 19 37 76 70 33 506 37 1 20 37 100 00 per cent.	(Analysed after dry- ing) 17 fb per cent. 2 03 " 10 83 " 10 94 " 2 27 "
Matter solubje m water,	21 58 per cent	29-28 per cent	24 85 per cent

R Wolffenstein (Zeitsch. augsw. Chemie, 1890, No 3) has described two varieties of factitious coffee, respectively known in Germany as Domkoffee and Allerwelthoffes. Both preparations were calriedy destinate of caffinne One consisted practically of a 11 o or y, while the other contained large quantities of 1 up 11 n e s. From the latter specimen Wolffenstein isolated a brown colouring matter having the specinescopic and chemical characters of Cassella-Drown. It was soluble in alkalies and in wister, but was completely precipitated from its solutions by hydrochloric acid. Fourteen grammes of the sample extracted with water and pre-cipitated with value of 10 for gramme of the colouring matter (1):

Foctious coffee-bean are, with very rare exceptions, heavier than water, while genume rousted beam are invariably lighter, unless much over-rossted. In taking the specific gravity, twenty beams should be immersed in brine, which is then diluted with water till ten of the beams float and the remainder such. The result shows the average density, but individual factitious beams often vary considerably from the mean.

In genuine coffee-beans a portion of the fine membrane or "parchiment" with which the bernes were invested will almost always be found adhering in the cleft. The microscopic structure of the bean, as seen in a thin section, or of the powder affords a certain means of recognising its nature. Most factitious beans contain starch, which is entirely absent from gaunus coffice Cincory and other roots are readily recognisable by the microscope. The methods used for the examination of ground coffee may also be availed.

D'an gway bean as, the seeds of Cassat tora or O cocidentality shundant an Entain Burmah, have been prepared and patentied as a substitute for coffee (Eng. Pat., 15,564, 1888) In Germany, the ground and rosated seeds have been sold under the name of "Mogdad coffee," and it is said that a smalles proposition than 20 per cent. in coffee cannot be detected either by the taste or the appearance of the sample. Dangway beans leave about 10 per cent. or said on ignition, and have a charactasistic moncopic appearance which has been described and illustrated by A. Wyn iter B 1yth (Food, Composition and Analysis) They sink very rapidly in watea and colour brine more intensely than do coffee beans Dangway beans contains a famin distant from caffeianme and. They are destitute of caffeine, but O. Hehn er has detected a minute quantity of some other alkaloud of his relation of the content of

The use of Mussaenda Borbonica seeds, to be mixed and reasted with coffee-beans or entirely substituted for them, has also been patented (Eng Pat, 14,945, 1888) 1

¹ Investigations at Kew Gardens show the supposed Mussanda seeds to be really those of Gartnera vaginata. They contain no caffeine.

The beams of a species of Phassoius are reported by E Frick e to be reasted, ground, and sold as "Congo cofice" The berries are very large—214 filling a 100 cc mcseure—and of shining black colour The infusion is very sattingent and contains no seffunce or their cavatilisable alkaloid.

To distingual hupne-seeds from coffee-beans, Hager texts 3 grammes of the powdered sample with 20 cc of water and filters after half an hou. The filtrate from genume coffee will be foelyly qellow and not taken in the least degree bitter, while in the presence of lupne-seeds a marked bitter taste will be observed.

GROUND COFFEE Beaules the foregoing sophistacians and substitutions of the coffee-bean ground coffee is liable to various adulterations. Some of these can be telerated when practised in moderation, provided that the fact and proportion of admixture and dily acknowledged; but it mast be remembered that all these additions, inclinding chaory, the least objectionable and by far the most widely used, are destitute of the volatile out and peculiar alkaloid which give to coffee its most valued properties. The dimminished consumption of offse in England is doubtless largely due to the frequency and extent of its sophistications.

¹ The late Dr Wm Wallnes, writing in 1886 (Assipit, ix 42), names the following articles as used for shittening offer—Chicory, seamed, from the forested figs, dired dates, date-stone, decayed ship basents, beans, possessing, milt, dandelson toot, tuning, cariots, parsuege, and mangield-warned Damaged runsins are stated by Albert Smith to be used together with choosy for unking Premet onfer.

*CONTINUED CRITICAL IN PRESENT FROM the 100 of Conformer malphus, which is not thin obsee, blind-side, and then issaid in this same manus at coffse, usually with the additions of a small proportion of fat of some kind for preparation and use of reasted choory appears to have organized in Holland about 1750. A Mayer (Red Contral, 1855, page 328) gives the Hollowing as the compaction of these samples of Dick honcy not — Water, 7700 to 77 3 per cent , albummonds, 11 , fat, 02, invilue and other ronincegoness matters massable in analond, 120 to 17 3, cando filter, 140 to 18, sagar, &c., 6 00 to 60, bittle extract, 0.05 to 0.15, and sah, 140 to 19, control Mayer from the batter substances strated by chierofetim to be calculated in water and alcohol, unsofulled in which and also bed by conducted the control of the control

A Peter mann (Bud. Central, 1883, page 843) gives the following results of analyses of two samples of roasted choory, one of which was consely and the other finely ground. The sah was somewhat higher than usual, but was perfectly white. The fat shown was probably not all natural to the The chief adulterations likely to be met with in ground coffee are.—(1) Mineral matters, (2) roots, such as chicory, dandelion, turnip, (3) seeds and seed-products, such as beans, aconas, and cereals, and (4) saccharine matters, such as caramel and reasted dates and figs.

In Bulletes No. 29 of the Laborstory of the Inland Rovenue Department, Cannad, the chart analyst, T Mocfarlane, states that——"There are, uncrover, large quantities of a substance unported under the name of eace are of coffee, for adulerating purposes, which is a species of humi sugar, and, from its containing dixtrin, is probably made from some of the byperochoics of the glucose factories. It code in New York and Philadelphia from 3 to 15 cents per 16 As it possesses no organic attachart it as pit to even for the containing power, and a little of it is capable of imperting a atong bown coffee colour to water."

Caramel, when added as such, may often be distinguished under a low microscopic power by the jet-black colour of the particles These dissolve easily in water with intense brown colour, and the solution has a lutter teste

A factitious caramel is now manufactured by adding to glucose about one-eighth of its weight of a brown coal-tar dye, naphthol-brown.

A useful prelimmary test for ground coffee consists in goulty stewing some of the powder on the surface of cold water. The oil contained in coffee prevents the particles from being readily wetted by the water, thus causing them to float. Chicory and the chicory, as the proportion recorded is largely in excess of that found by other observer. The water also is much above the usual proportion in recording the control of the

	Coarse Grains.	Fra	Powder
Water (lost at 160°-105° C),	 16 28	16.96	
Gincowe,	28 12	28 70	
Dextrin, inulin,	9 63	9 31	Soluble in
Albuminolds	8-23	8 65	hot water
Colouring matter and bitter extractive,	16 40	17 59	
Ash in soluble portion.	2.58	2.55	
Ash in insoluble portion,	4 58	5 89	
Albuminoids.	15	2.98	Insoluble in
Fat,	571	3-92	=26 11
Cellulose,	12 32	18 37	,

majority of coffee adulterants contain no oil, and there examels as every quantity extended by the water, with production of a leaver oolone, while the particles themselves rapidly sink to a leaver older, while the particles themselves rapidly sink to the botom of the water. To strain he house, offee becomes tolerably uniformly diffused without sensibly colouring the water, while choory and other sweet roots quackly give a dark brown, tarbid unfusion. Roasted careles do not give so distinct a colour.

According to A. Frans (Arch. Pharm, [5], vr. 298), if 2 c.c. of a 10 pat cent infusion of coffice in boiling water be treated with 08 c of a 2½ per cent, solution of cupric sectate, and the liquid filtered, a greensh-yellow filtrate is obtained. If chiccey be similarly treated, a dark red-brown filtrate results, the colour of which changes on standing Ten per cent of the adultorant can thus be detected.

The colour of an infusion of chicory is said to remain unaltered on addition of a solution of farme oliurida or sulphiats, while the brown colouring matter of coffee infusion turns groun, and is pairably prespitated as blush-green flakes. In am infusion of mixed chicory and coffee, the reagent forms a precipitate, and leaves—though more of the prespitate is facilitated by tendering the liquid alightly alkaline by ammonia (Dirach's solut Jour, exit / 78).

Albert Smith (Pharm Jour, 13] xz. 569) resommends, for the detection of checory in office, that 10 grammes of the sample should be boiled with 250 c c of water, and the liquid strained and precipitated with a slight excess of base leed accetate. On allowing the precipitate to estile, the supernatant liquid will be colouries it pure coffice has been undoor teatment, but in presence of chacory will be coloured to a groater or less degree according to the proportion present, which can be estimated from the depth of tint by a process similar to that of nesslerising water.

The three foregoing tests are occasionally of service for the examination of infusion of coffee when the solid article is not available, but they cannot be regarded as so estisfactory as the actual recognition of the adulterant by the microscope.

The great majority of seeds likely to be met with in coffee contain a notable quantity of starch. This is true of hean, peea, acouns, and all cereals and products therefrom Hence if starch be absent, the freedom of the coffee from all this class of adultorants is certain If present, the nature of the admixture can usually

If a funnel be used for the above test, the sunken particles may be readily let out and examined under the misioscope. be ascertained by a microscopic examination of the prepared sample.¹

For the detection of stamb, the author boils the coffee for a for numtes with about 10 pairs of water When the luquid has become perfectly cold, some did to suphurue and is added, and then a strong solution of permanganate of polassum diopolar cautionally, with agritation, till the colouring matter is neadly destroyed, when the luquid is straned or decented from the soluble soluble matter. On now adding a solution of solume to the solution, a blue colourion will be produced if any startly solution.

Some operators employ animal charcoal for decolorising the coffee infusion before testing for starch. The addition of starch-holding adulterants to coffee, in the author's experience, is rate, but in the United States and Canada is very common, the adulterants there found including wheat-flour and bran, buck-wheat, barley, maize, peas, peas-hulls, &c **

The insoluble matter remaining after treating the coffse with water and decolorising with permanganate can be advantageously examined under the microscope for chicory and other non-starely additions, the structure of which is more readily observed after the romoval of the colouring matter.

F. M. Rimmington (Pharms, Jour. [3], xz 599) recommends, for the removal of coloning matter, that the sample of coffee should be boiled for a short time with water containing a little enrolonate of sodium: After subsidence, the liquid is poured off, the results washed with water, and then treated with a weak solution of bleaching powder until decolonastion is effected, which usually occurs in two or three hours. The real coffee will then form a dark stratum at the bottom of the beaker, and the choory a light and almost white stratum floating above it, and showing a clear and sharp line of separation.

- ¹ For this purpose the coffee should first be exhausted with ether to remove fat, and then treated with methylated spirit to dissolve the colouring matter. In the residue, the starch and other structures will be readily perceptible.
- A certain famous sample of coffee alleged to contain acoins gave the author no reaction by the above test, but after the addition of 2 per cont of reasted acords the test showed the presence of statch very clearly
- In 1876 a large searnte was made in the east of London of a mixture of until 90 of loasted account Reasted acoust were first placed before the English pubble as "Pelotas coffice," and arbsequently as "coffee surrogate," but the manufacture of both these preparations was stopped by the excess.

Under the nucroscope, chrony is readily recognised by the pseudiar dotted appearance of the vessels, often occurred in bundles, and by the characteristic appearance of the large cells. Dandelion, turnips, and other sweet roots present a close similarity to chroory, and can only be sately destanguished therefrom by careful microscopic comparison of the sample with the actual roots in question.

The microscopic appearance affords the only certain means of identifying choory and other roots in coffee, and the same statement applies to saccharine fruits, such as roa-ted figs, dates, raisins. &c. 1

The nature of an adulterant of coffee having been assertationd by the and of the microscope or other means, an attempt may be made to deduce the proportion present from the olumical compontion of the sample. When only one adulterant is present, this may sometimes be effected with a far approximation to accuracy, but even in the case of clusory it is not always possible to ascertain the proportion within a somewhat wide limit.

For ascertaining the proportions of adulterants in coffee, the only chemical distinctions of any practical value are —Certain constituents of the ash; the proportion of fat as extracted by ether or petroleum spirit; the proportion of aque ous extracts.

³ Printed descriptions of nucroscopic characters are of little value, and drawings are often muleading. The adulterants of coffee are best examined as transparent objects under a moderate power, and, except where statch is to be identified, by unpolarised light.

² What can be done in this manner, and the errors hable to occur in practice with deficient methods or imperiction ampliation; is apparent from the following figures obtained in 1882 by various analysis to whom exactly similar asymptotic of mixed ceffee and chicary of known composition were submitted (Analysi, vii 76).

Actual percentage of Chicory in sample,	10 per cent	30 per cent.	374 per cent		
Percentage of Chicory reported Somerset House (Referees),	Not more than 21 per cent	Not less than 25 per cont	Not less than 43		
Δ,	7 per cent.	31 per cent	85 per cent.		
В, .	7 ,,	32 ,	34 ,,		
О, .	5 to 10 per cent.	25 ,,	50 ,,		
D, .	16 per cent	86 ,,	47 ,,		
В,	Genume	81 ,,	50		
r , ,	Upwards of 10 per cent	Upwards of 80 per cent	Upwards of 40 per cent		

as deduced from its weight or the specific gravity of the solution, the colour of the infusion, and the proportion of caffeine in the sample. In all cases of importance two or more of these methods should be applicable.

A. Smetham (Analyst, vn. 73) obtained the following range of figures by the analysis of seven samples of roasted coffee, representing typical commercial qualities.—

Moisture (lo	at at 10	00° O),			1 59	to	3 .89	per ceu
Oil (other ex	ktract),				10 18	,,	12 13	- ,,
Caude fibre,	1,				70 84	,,	74 50	,,
,, ,,	m sam	ple dne	d at 1	00°, .	78 71	,,	75 70	,,
Cellulose,					28 84	,,	84 40	,,
Nitrogen,					2 14	,,	2 38	,,
Total ash,					4 08	,,	4 63	,
Soluble ash,					8.14	,,	8 60	,,
Ratio of tot	al ash t	o solubl	а		100 72		100	82 ,,

The following analyses by C Krauth (Ber, xi 277, Josep. Chem Soc, xxxv. 449) give some comparative figures for coffee and its more probable additionants. Except in the case of the last column, the results apply to the substances previously dried at 100°—

			Sug	ar			Moisture	
	Ash	Fat	Pre- existent	After Boiling with Acid	Soluble In Water	fn	in Undried Substance	
Coffee, reasted, five	4 19 to 6 88	11 78 to 15 6	} 0-2	24 29 {	22 47 to 25 21	74 70 to 77 58	1 47 to 4 87	
Chicory, ressted, .	10 83	1 15	23 40	22 14	85 42	34 58	4 80	
Chicory, unrossted,	5 85	48	23 84	Not do-	7871	21 28	6.89	
Rye, rosated,	2 48	1 68		75 37	31.92	68 07	0.28	
Wheat, rossted,	1 80	2 75			52 65	47 85	1	
Coffee, with 10 per }	4 81	14 16	19	29 66	25 98	74 46	2 15	
Coffee, with 10 per }	5 10	12 55	2 80	23 15	80 68	69 86	2 30	

¹ The "crude fibre" was determined by boiling 2 grammes of the sample with three successive quantities of water, and washing the residue on a counterpoised filter till the washings were colourless, when it was dried at 100° C and weighed

The following analyses by Konig show the composition of certain adulterants of coffee

	Ohleory	Pign	Acorns	Rye
Water,	12 16	18 98	12 85	15 22
Nitrogenoue matters,	6 09	4*25	6 13	11 84
Pat,	2 06	2.33	4 61	846
Bugar,	15 87	34 19	8 05	,803
Other non nitrogenous matters, .	48 71	29 15	82 0	55 87
Celluloso,	11 00	7:10	4 98	5 35
Ath,	619	8 44	` E-19	4 81
Matters soluble in water,	68 05	78 81		45 11 (

The following table shows the published results of analyses of coffee substitutes and to be manufactured respectively from access, i.e., and barley 1—

								"Acom1 Coffee"	"Ryo Coffee Bubstliute "	"Rarley Coffee	"Barley Coffee."
Water,	_	•		,				12-85	2 22	9:45	041
Nitrogeno	as I	matt	678,					618	11 87	9 38	10 58
Fat,								4 01	8 91	3 25	1.04
Sugar,								8 01		1	
Starch,)	8 84		
Daxtelu,								62:00	49-51	70 13	08 88
Other non	nit	roge	nou	me	tten	и,)	88.9)	
CeUulose,								4.98	9-78	4 26	10 50
Ath,								2 02	6 54	8 36	3'04
Matters so	lub	le in	wai	ter,					61.83	31 20	84'37
Glucose f	orn	ned turio	by acid	boi 1,	ling	wi •	th	}		60 28	07 19

Moscheles and Stelzer have recently published complete analyses of several coffee substitutes (Ohem. Zeit., 1892, xv. 281; Analyst, xv. 154). One of these contained hipmes (which they consider a very reprehensable addition), and enother was destitute

¹ The "courn coffee" was analyzed by König, who found from 20 to 30 per cent of starch, and 6 to 8 per cent, of a variety of taunus acid. The "rye coffee substitute" was prepared by Bohr Bros The analyzes of "barley coffee" are by C Kornanth.

of coffee, but contained 0.31 per cent of caffeme, due to the presence of nowdered kola-nut.

The cash of pure coffee is generally between 3½ and 4½ per cent, rawly, if ever, exceeding 6 per cent, and even when a considerable proportion of choory is present it seldom, rases beyond 6 per cent. Any notably higher proportion will indicate the presence of a mineral adulterant. The sah should be white, or nearly so, any marked ted this undicated an added compound of non.

The composition of the sah of coffee presents some marked differences from that of chicory, as is apparent from the following results of analyses by H Ludwig (Arch Pharm, [3], 1, 482) and James Bell (Foods, 11, 46, 57).

-						Coffee H L	beans udwig	Ooffee-beans, Eight Samplos	Ohicory Root, Eight Samples J Bell			
-						Gness Soll	Timestone Soil.	J Bell.	Deducting SIO ₂ and Sand	Including 8:0 ₂ and Sand		
1	K ₂ O	_		-		14 13	44 03	58 20 to 55 80	27 85 to 46 27	24 88 to 83 88		
1	Na_20					5 84	5 85	Not detected	8 17 ,, 10 90	2 04 ,, 16 10		
Ì	CsO					8 64	4 89	410 to 818	7 65 ,, 10 81	500, 900		
į	M_00					814	8 01	820 ,, 887	588 ,, 806	3 42 ,, 7 22		
-	Fe ₂ O ₃					18 54	198	044 ,, 098	3 50 ,, 8 20	3 18 ,, 5 32		
ì	P_2O_5					18 65	10 54	10 15 ,, 11 60	9 50 ,, 12 61	6 85 ,, 11 27		
money	803					15 28	1 64	809,, 520	8 88 ,, 11 78	5 88 ,, 10 58		
-	O1 .					Tinos	0.98	0 28 ,, 1 11	508, 608	8 28 ,, 4 98		
I	σο ₉ .					8 84	21 24	14 92 ,, 18 13	2 04 ,, 4 60	178 ,, 819		
-	2018					1 65	0 37	0.00 ,, 0.45		2 61 ,, 12 75		
-	Band					None	None	None		8 08 ,, 23 10		

Ludwig found in each case a notable amount of sods, a result which disproves Ball's improbable statement that this base is absent from coffee-ash Ludwigs figures also show an enounous variation in the proportions of $K_{\rm L}Q$, $E_{\rm S}Q_{\rm S}$, $E_{\rm L}Q_{\rm S}$, and $CQ_{\rm S}$ according to the nature of the, soil on which the ordize-plant is grown 'If the Na₂Q in chicacy-sub be calculated into its semivalent of $K_{\rm L}Q$, and the figure thus found added to the actual $K_{\rm L}Q$, the percentage is not greatly different from the proportion of potent found

The sample of coffee from a gress soil must be regarded as highly abnormal. In the wide experience of the author the sah from genume coffee has never been observed to have a red colour, as would be the case with the ash of a

by Bell in coffee-sals. The proportion of exide of iron is notably greater in chercy than in coffee. Hence cluczy-sals always has a red tange which is absent from the ash of genuine coffee. A notable difference is observable in the proportions of CO₂ and Cl, and a very wide distinction in the figures for sand and sitica. In only one of the eight samples of coffee did the situa even approach of per cent, and in another pottom of the same coffee, which was properly sercenced before lossing, the silica of the ash fell to sail.

In consequence of the large proportion of potassium carbonate in coffee-ash, the porcentage of the total ash soluble in water is much greater than in the case of chicory-ash, and attempts have been made to utilise this fact for ascertaining the proportion of chicory present in mixtures of the two. Thus the author found from 60 to 85 per cent of the total ash of coffee to be soluble in water, whereas on an average only 34 per cent of the total ash of chicory was soluble in water. But this proportion is gravely affected by the proportion of actual sand which may be present. This values in commercial chicory from a trace up to 4 5 per cent. which difference is quite sufficient to invalidate deductions based on the ratio of the total to the soluble ash. By comparing the soluble ash with the total ash minus sand and silica, somewhat more rehable results are obtained, but at best the method is only capable of affording a rough indication of the proportion of chicory present It may, however, serve to point to the presence of a foreign ingredient, which can then be identified and determined by other means The following ash-analyses, by James Bell, are interesting in this connection -

	Lupina	Acorna	Mairo	Parentps	Dandelson Root	
K20	33 54	54 98	30 74	66 54	17 95	
Na ₂ O	17 75	0.68	Not found	Not found	80 95	
CnO	7.75	0.01	8 00	6.85	11:43	
MgO .	618	4 32	1472	0.49	1 31	
l'egO ₃	ì	0.54	0.84	0.63	1.27	
PgO5	25 53	11 16	44 50	18 84	11-21	
80 ₃	6 80	4 70	4 13	4 07	2 87	
CI	2 11	2 51	0 50	2.00	3 84	
CO2	0.66	18-69	1	11 44	6 21	
SiO ₂ , &c	0 87	101	178	0 57	11 26	
	101 09	90 58	100 27	102 42	97 80	

The following centesimal figures by Way and Ogston refer to the ash of other roots:

	Turnip	Beet	Carrot		
TegO2	0 14 to 0 08	0 52 to 3 74	0 59 to 1 65		
cı .	8 ,, 5	85 ,, 30	8,,46		
002 .	95,, 15	15 ,, 21 6	15 ,, 19		

The fat of coffee is tolerably constant in amount, and hence the proportion services as a useful undication of the amount of certain admixtures. The owns Macfarlane, Head Chemist of the Inland Revenue Dopartment, Othawa, informs the author that the petroleum-ether extract from previously dried coffee langes from 10 to 12 per cent. Only one sample out of nearly fifty examined showed less than 10, and no sample gave as much as 18 per cent, which is previously interesting the contract of the

The aqueous extract of coffee is remarkably constant in amount, and in very little affected by variations in the neasting. Instead of weighing the actual extract, G raham, Stenhouse and Campbell (Journ Chem Soc, Ix 88) determined the specific gravity of the aqueous infusors of coffee and various roussed vectable matters. Their method was to treat the rosstell substance with ten times its weight of cold water, mass the liquid of the boiling-point, and observe the density of the filtered liquid after cooling to 60° F. (= 15 6° C.). The following is a classified arrangement of their results.

Substance	Specific Gravity of 10 per cent Infusion	Substance	Specific Gravity of 10 per cent Infusion
COPPEE — Macha, Mulgherry, Plantation Ceylon, Java, Janusce, Native Coylon, Oosta Rica, Costa Rica, Average, Lacournous Sends— Lanins	1008 0 1008 4 1008 7 1008 7 1008 8 1009 0 1009 0 1009 0 1008 7	Roors — Chilory, Yerkshire, Begish, Poreign, Guenvey, Averago, Carroba, Turrips, Dandehen, Red beek, Mangold wurzal,	1019 1 1021 7 1022 0 1022 0 1023 0 1021 05 1014 3 1017 1 1021 4 1021 9 1022 1 1028 5
Pens, Beaus, MISCELLANEOUS -	1007 8 1008 4	OBBEAL PRODUCES Brown malt, Black malt, Rye meal,	1010 9 2021 2 1021 6 1025 3
Spent tan, Acorns,	1002 1 1007 8	Maize, Bread raspings,	1020 3

These results show a marked distinction between coffee, leguumous seeds, and access on the one hand, and cereal products and cheory and other roots on the other Unfortunately, with the exception of cheory and coffee, they apply merely to single specimens of each kind of substance

Experiments made in the author's laboratory gave a mean density for collect-influence pressely identical with that obtained by Gruham, Stanhouse and Campboll (1008 7) Operating as they prescribe, however, there is a dways a suspensor that the exhauster is monuplete, especially in the case of genuine cofloe which has not been very finely ground. Hence in a series of experiment made, in the author's laboratory, the sample of coffee was well bouled with 10 parts of wiser, the solution filtered, and the centime washed with hot water till the filtrate measured 10 cc for every 12 gramme of the substants employed. Operating in this rannance, the influence from fourtheen specimens of ordinary commercial reaction offsee (ground in the laboratory) were found to have a specific gravity imaging from 1008 8 to 1008 5, with an average of 1007 9 1 (Asalost. v. 1)

J Skalweit has shown that the specific gravity of the aqueous infusion is not sensibly affected by the extent to which the coffee has been reasted

By the exhaustion-process, the author obtained the following results from samples of commercial chicory (undried) —

					٠		6	pecific Gravity I 10 per cent Infusion
Yorkshire	Chicory,	under-10	asted,					1025 9
	,,	(same san	mple), l	nghl	y roasi	ed.		1019.0
Chacory of	unknow	n origin,			٠.			1021'1
,,	,,							1020 0
**	**	19						1023 4
					Moan,			1021 9

It is evident that the density of elincory influsions varies much more than that of coffee, a fact which pievents the method from farmishing more than an approximate determination of the proportion of coffee and chicory in a mixture of the two. A shaper result may be obtained by pievrously drying the sample at 100°,

¹This figure is comewhat lower than the average of Guham, Stenhouse, and Campbell's experiments, which trads to above that they offered practically perfect exhauston. The differences is not improbably due to a slight ions by expectation when the influence is made by reasing the liquid to the boling-point, instead of boling thoroughly and making the influence approach in the order of boling the origin yand on making the influence approach is a first solution measure after solution. (I he is no lies neet with a genuine coffee giving an influence distinct of 1010 2.

and hence chimmating the somewhat senous error due to varying proportions of mosture Adopting 1024 as the normal gravity of the infusion of drace choicey and 1009 as that of dract collec, the percentage of real coffee in a mixture of the two will be found by the following equation, where d is the specific gravity of the 10 per cent. infusion and C the percentage of coffee in the sample:—

$0 = \frac{(1024 - d)100}{15}$

A McG-111 (Trans-Royal Soc Canada, 1887) finds that the cleanty of the minacons of colles and chaory is materally affected by the finances of the powder, the time occupied in beating the decoction to bothing, and the time during which the boiling with water is continued. He recommends that a weight corresponding to 10 grammes of the mousture-free sample should be boiled with 100 cc of dishiled water in a flast fitted with a refux condenser. The less is adjusted so that sbullino commences in no to fifteen munities, and the boiling is continued exactly one hour, when the flame is removed, and after fifteen number's rest the liquid is pressed through a dry filter. The average donaity of a 10 per cent election of pure coffee thus prepared was found to be 1009-85 at 52°, the mean number for choory decochom (three samples) being 1028-91 at the same temperature, or a difference of 18 30. From these results the following formula may be deduced:—

1 Thos. Macfarlane, Chef Analyst in the Inland Revenue Leboatord, by Chiwa, has communisated to the sulfror the Glowing results, oblation by the application of McGull's method for secretaring the infessor-density and costal determination of the soluble extract. This last determination was made by thoroughly extracting the dried sample with petroleum ether, and then treating the refracta ellestance with boding water. Instead of everpositing the adultion, the macluble matter was redried and weighed, the loss showing the "water extract".

						Water Extract	Infusion Gravity at 02° F
20 20 20 20 20 20 20 20 20 20 20 20 20 2	th 10 per , 20 , 30 , 40 , 50	cent	Chicory,	: ::	:	22 41 21 92 20 42 25 90 25 90 30 75 37 40 43 84 55 93 60 54 66 53 71 41	1009 78 1000 73 1011 68 1013 14 1015 28 1017 08 1018 08 1020 48 1022 70 1024 15 1028 42 1028 32

It is evident that the specific gravity of the especies infrason is really a function of the solid matter dissolved by the water, and a close approximation to the percentage of the latter can be obtained by dividing the difference between the solution-density and 1000 by the number 0375 or multiplying it by 2671 Thus if a coffee-infrason have a density of 1009 0, the proposition of matter solide in water will be

The figures for soluble extract obtained by T. Mn of or land (Ottawa) by the analysis of 64 samples of commercial order ungged from 215 to 265 per cent, with an average of about 24 per cent. The samples were duted at 100° , deprived of fat by treatment with pertoderm either, reweighed, and then exhausted with water. Instead of evaporating the infusion and weighing the soluble extract. In a scale of the control of the co

Alfred E. Johnson states the soluble extract from previously dried (reasted) coffee to be very constant at 24 pc; cent, and the extract from dried choosy to eventage 70 pc; cent, and on these figures bases the following process for the analysis of coffee mytures.

The ground coffee as dreaf at 100° C, and 6 grounness weight of the mosture-free sample belief of fifteen minutes with 200° c. of water. After settling for a few minutes, that injund is powered off though copies wine-gause or coarse mushin into a 250° ce flast. The grounds are boiled with 50° to of water for two minutes and the liquid strained as before. The contents of the flask are cooled, made up to 250° c. gatated, and power of the filtrate, rejecting the first portion (equal to I grainme of the dry sample), as then evaporated in a flat dath over boiling water,

¹ This factor is deduced from the known solution densities of caranel and the calbohydrates. J. Skalweit (Rep. Anal. Chem., 1882, page 227), as the result of direct experiment, gives the following data—

At 17 5° O., 1 001 sp. gr. of 20 % infusion represents 0 36 extract per 100 c.c.

⁹ O Hehner found a hightly-reasted cheory (drod) to give 67 1 per cent of soluble matter, and an infusion-density of 1024 4, while a highly-reasted sample had an infusion-density of 1019, and yielded only 54 1 per cent of extract.

^{. 1 235 , 48.25 , 18.25 , 18.25 , 19.2}

and the residue (representing the extract from 1 gramme) dried in the water-oven and weighed Then —

100 (70 - per cent of extract found) - percentage of ooffee in sample.

The results thus yielded by coffee and its principal adulterants are given on pages 543, 544

The tinctorial power of the infusion was suggested by Graham, Stenhouse and Campbell (Jour Chem. Soc, 1x 36) as a means of determining adulterants in coffee They found that the depth of colour of the liquid obtained by infusing coffee and its adulterants in 2000 times their weight of boiling water varied remarkably, caramel giving about seven times and chicory about three times as deep a colour as coffee 1 But their experiments showed that four different samples of pure coffee varied in tinctorial power between 143 and 183, as compared with caramel as 1000, and no doubt samples of chicory would be found to present at least as great difference in colouring power, according as they happened to be lightly or strongly roasted Nevertheless the author found (Chem News, xxix 140) that the tinctorial power of an infusion of mixed samples of chicory was almost exactly three times that of an infusion of average or mixed coffee, and that different samples of chicory did not vary more than from 2 8 to 3 2 m colouring power when compared with the same sample of coffee estimate the proportion of chicory in a sample of coffee mixture, a standard mixture should be prepared by mixing together several representative samples of genuine ground coffee with an equal weight of mixed chicory 2 One gramme of this standard coffee mixture (containing 50 per cent of coffee), and the same weight of the sample to be tested, are boiled for a few minutes with 20 cc

¹ The following are the relative amounts of various reasted substances found by Graham, Stenhouse, and Campbell to impart an equal depth of coloni to the infusion —

2 50 | Coffee, 5 2 85 | White limits seed, 1 00 | Paranips, Caramel. . 10.00 Mangold wurzel. 1 66 Maize and rye, 8 43 Beans and peas, 1 82 Dandelion root, Black malt. 8 33 Spent tan, 83.00 2 00 Red beet, White turnips. 2 00 Bread tasnings, 8 64 Brown malt, Committee 5 00 Chicory (darkest Yorks), 2 22 | Acorns,

It the standard coffee mutton be kept, it under goes a change which modifies, even use day teach, the colours of the influsion A. persuasent steadard of interest of the influsion A. persuasent steadard of interest of the influence of the observation of custom in stead can also be employed as a standard, if it at waits to personally ascertained that that are best observed by placing a piece of wet filter paper behind the tobes while they are held up to the light?

of water. The legada are cooled and passed through a double little, the moshible portions being repeatedly belief with fisch quantities of water till no more colour is extracted. The solution of the standard mixture is then made up with water to 200 e.c., and the solution of the sample to 100 e.c. Ten e.c. of this latter liquid is poured into a narrow graduated tube, and some of the standard solution into another tube of excelly equal bore. If the sample convists of pure coffee, the two liquids will now be of excelly smulier into; but if cheepy be present, the solution of the sample will be the darken, in which case water is gradually added hill the tints are precosely equal. When this point is attained, the volume of the sample solution is observed. Every 1 e.c. of water added represents 5 per cent of clucery in the sample. Thus I the liquid measure 17 e.c., the sample contains 55 per cent of cheery,

J. R. Leebody (Chem Nows, xxx 243) has described a similar method, but, instead of observing the colour of the solutions transversely, he dilutes the solution from 1 graume of the coffee to 700 cc and observes the colour from above, as in nesslerising water.

The observation of the infusion-colour is occasionally very useful as an indication of the presence of caramel added as such, since in that case the colour will be greatly in excess of the proportion of chicory or other adulterant as deduced by other methods.

The caffeins of coffce is tolerably constant in amount, and hence its determination has been recommended by Paul and Cownley (Pharm Jour., [3], xvii 565, 648, 821, 921) as means of estimating the proportion of real coffee in a mixture. These chemists have shown (page 492) that most of the published methods for the determination of caffeing give results more or less below the truth. but that when the process recommended by them is adopted the proportion of caffeine isolated varies within comparatively narrow limits. This is especially the case if the roasted berries are dried at 100° before grinding thom, as by this means the error due to variable proportions of water is eliminated, and the coffee can be obtained in a finer state of division, and hence be more perfectly exhausted In fourteen commercial samples of coffee-herries, Paul and Cownley found the moisture to vary from 6.2 to 100 per cent. After drying at 100° C, the caffeine ranged from 1.20 (in a coffee . from Coorg) to 1.29 per cent (found in coffee from several sources), except in Liberian coffee, which yielded 139 per cent. On the basis of 1 3 per cent of caffcine in genume coffee, adopted by Paul and Cownley, the proportion of real coffce in a mixture will be found by dividing the percentage of alkaloid found into 130. It would be safer to adopt the number 120 instead of 130, and in using the method great care is necessary to effect the isolation of

the whole of the cafform. To ensure this, the sample must be in very fine powder, the exhaustion by alcohol of the mixture of coffee with lime or magnessa must be proceed to be complete, and the egitation of the autosous input with chlorotorm must be repeated until no more alkalod as exhauted.

Although, when taken alone, any one of the foregoing methods of examining coffee is hable to lead to determinations of the proportion of adulterants somewhat wide of the truth, by the combined use of several a fairly accurate deduction can be made In certain zero cases, additional information may be obtained from the determination of the fatty matters, the alkalimity of the soluble ash, and the proportion of mitrogen.

Corress Extraors are prepared with very limited success by subjecting roasted coffee to treatment with boiling water or steam, and adding the volatile products to the squeous extract. The product is deficient in saffeine, and does not constain all the extractive matter of the coffee, now, when dituded with the appropriate amount of water, is the colour the same as that of the treshly-prepared input. To remedy this defect caramel is added, together with strong alcohol as a preservative. In one patent, addition of chorcy and singue is preserved. The following results were obtained by A D omergue by the examination of six samples of coffee extract.

Î			Water	Extract disod at 100° U	Caffeino.	Ash		
ļ	Α,		,		88.8	13.7 per cent	0 108 per cent	0 61 per cent.
١	В,				824	176 ,,	0 105 ,,	0 70 ,,
	O,				58 99	41 01 ,,	0.080 **	4 30 ,,
ì	D,				72 8	27 2 .,	0 040	8 10 ,
-	Ð,				8 98	301 ,,	0.060 ,,	140 ,,
1	F,				80 74	19 28 ,,	g 000 ,,	,183 ,,

Samples A and B were prepared in the laboratory C, D, and E were coloured with caramel Domergue regards the proportion of caffeine as the best indication of the value of a coffee extract.

Of three samples of "coffee extract" examined by G. L. Spencer, one was destrate of caffeine, but contained corasion and other starchy bolies, a second contained 119 per cent. of caffeine, or about as much as ordinary coffee, and a third was a maximum of coffee extract with milk and sugar, and contained 072 per cent of caffeine. Very notable proportions of im and copper was described in these proparations

Kola-nuts.1

The G our ou or Kolanut, from a tree belonging to the family Sterouliacea, is chewed and used for preparing a beverage in Western Africa, by the negro inhabitants of the West Indies, Brazil, &c

From the nut of Sterculsa or Cola acumunata, the female or true Kola, Heckel and Schlagden hauffen (Pharm. Jour, [3], xiv 584) obtained the following products.—

(Caffeine,			2 348	per cent.
Extracted by	Theobrom	me,		0 023	,,,
Chloroform -	Fats.			0 585	. ,,
ł.	Tannın,			0.027	
ì	Tannin,			1 591	**
Extracted by	Kola red,			1 291	
Alcohol -	Glucose,			2875	**
1	Salts,			0 070	
ì	Starch.			33 754	
i i	Gum,			3 040	
1	Colouing	matter	s,	2 561	11
Undissolved	Protends,		•	6 761	
ì	Cellulose,			29 830	
	Ash,			3 3 2 5	,,
į	Water,			11919	<i>"</i>

According to E. K. n. b. b. i. (Apoth. Zett., 1899., p. 112), loadnuts contain a glucowde, ko l. an n., which no buding with water, or by treatment with dilute acults, sphits up into c. if fo in a, gluoses, and koln-red. (AgMajOM). This last products is an extremely unstable substance, taking up oxygen during the drying of the intis, with sepanation of water and foundation of gallotannic acid. (AgMajOM) its stated that fresh kolmuts have a greater physiological activity than when dired, as in the former condition the kolamin has not undergene the degeneration which destroys it and renders the caffein modibile

Monaron and Perrone state that powder and extract of kola-nuts have a far greater power of dimun-hing the elimination of phosphates and introgen than cassened alone has Kola-red has

Various other Afroan plants yield seeds closely resembling the true Kola, but it is doubtful whether they contain callenge.

¹ Kola-units are oblong, three forming a ball fully 2 inches in diameter, and resembling a very large berse-chestaut. The individual nuts have a rugged, dark brown surface. Insale they are hight brown, becoming risty on exposure, and tough as wood scirmont, and finally bittee.
After drying the bitterness dimminishes.

GUARANA. 555

a diminishing influence, but both it and caffeine act better in their natural combination than separately Caffeine has a diuretic action, whereas kola is anuretic. The drug prevents waste of brain as well as of muscular tissue.

FAINE KOLA, MAIR KOLA, Or KOLA BYTER, is the seed of Gracines Alota, a plant of the family of the Guttlerge growing in Libera and Contral Africa. On extracting the seeds with chloroform, either, and alcohol, no enfines to obtained, but only resms. One of these gives a violet coloration with fenne alls, while the other is destro-extency and preepintated by furtar emetic and bane lead acotate. The physiological action of the extract of kola latter is attituitiable to those receives.

Guarana.1

This product occurs in the form of cylinders. It is an undefinite mixture of various materials, of which the seeds of Paulisma sorbits appear to be the only constant and characteristic ingredient it is prepared by the Gu ar a n. a, a tribe of hall-savage Indians on the Upper Amazon. Its only interest is as a source of cassine, of which it contains a notable proportion. S ten ho use obtained 504, and F V G re on 505 per cent. E R S qui bb found 48 sper cent. Eghemens, n. 1615. J II F seems to C (Pharm Jour, 13), xm 363) obtained from 39 to 50 per cent of cassine to the control of the second of the control of the co

Cocoa and Chocolate.

Cocoa is the seed of the tree Theobroma cacae and allied species growing will in tropical America. It is cultivated in Branil, Grenada, Trunidad, &e., and has been introduced into the East Indies and parts of Africa and Aushaha. The cocoa-seeds from different districts vary considerably in appearance and flavour, but do not present any sharp distinctions in chemical composition

The fruit of the occa contains from 25 to 40 seeds closely packed in the pulp, which is removed by subjecting the seeds to a process of fermentation for a few days. The pulp is then separated by hand, and the seeds placed in trays and dried slowly in the sun or by artificial heat, boing turned over an intervals. The flavour

¹ Throughout Brazil, and m all parts of South America where the preparation 19 used, the word guarana 19 universally accented on the last syllable, and never monounced guarana

of the cocca is greatly dependent on the care and skill with which the openations of fementation and drying are conducted. The process has been compared to the maining of burley, germination taking places and being subsequently arrested. It is alleged that the alkaloid is formed during the process of fermentation, but the statement recurres confirmation.

When quite dry, the scoos-seeds are ready for exportation, but before bong used they are subjected to a gentle reading, whereby the butter tasts is modified and the keenels are more readily separated from the shells or busks, which constitute from 8 to 14 per cent, of the entire seed. When separated from the husts the broken kernels are known as occosion to 8.

Konig has published analyses of eight samples of decorticated cocoa-beans and of the hushs from the same specimens. The following figures show his average results ---

,	Monsture	Nitrogenous Matters	Fat	Starch	Cellu- lose	Ash
Cocca beans freed from	3 25	14 76	40 00	13 81	3 68	3 65
Cocca husks,	7 83	14 20	0.38	,	14 09	7:12

The following analyses of raw cocoa are by Boussingault (Ann Chin. Phys., [5], xxvm 433):--

						Kennel.	Kornel	Husk.
Water, Theobomine, Albuminoida, Asparagiu, Fat, Soluble collulose, Starch and glucose, Gunn, Tartaria said, Tamin, Ash, Undetermined.	:	:		:		7.0 32 109 truce 499 106 24 24 34 0.2	11 0 24 129 530 } 01 } 07	12 18† 14 25 3 9 12 12 2 5 05 6 89

[•] The presence of testaric acid in occoa has been confirmed by Wolgmann, who found from 4 St to 533 per cast in the raw whole bears. To determine it, he neutralised the squeeze extract with dimmons, added acidizing oblivide, redissolved the preceiptate in hydrochloic acid, and especipitated with socia.
7 This proportion of water scenes funcebably high

The author inclines to the opurion that the alkaloid of tea is in great measure a product of the decomposition of some more complex body, as has been proved to be the case with the cafferne of cola-nuts. It appears not improbable that the same may be two of the theoretismum of the coon-bean.

According to A H Church (Foods, page 200), good cocoamils contain —Water, 50, albuminoids, 170; fat, 510, theobromme, 15, cocoa-red, 30, gum, &c., 109, cellulose and lignose, 30, and mineral matter, 36 per cent

J. Bell gives the following as the composition of raw Trundad cocoa-nite.—Monsture, 5 23, fat, 50 44, starch, 4 20, alkalods, 0 84, albummous matters, soluble, 6 30, menible, 6 96; astrangent principle, 6 71; occoa-red, 2 20, gum, 2 17, cellulose, 6 40, medinite insoluble organic matter, 5 80, and ash, 2 75 per cent.

The following analyses of commercial raw cocoa, after removal of the husk, 'are by Eastes and Terry (Pharm Jour, [3], xy 764).—

Kind of Cocos,	Moisture	Fat.	Theo bromine	Ash	H ₂ PO ₄ ,
Caracess, Carapano, Gronada, Gronada, Grayaquil, Para, Surinam, Trinidad (common), Trinidad (fine, 38 Antonio),	4 75	58 65	1 08	2 70	1 88
	5 04	47 86	6 87	8 69	1 89
	5 50	47 12	1 42	2 81	0 01
	3 68	52 57	1 74	3 28	0 85
	4 80	57 67	1 60	3 09	1 88
	2 55	58 76	1 42	2 44	0 86
	5 62	45 71	1 05	2 70	0 89
	4 72	53 57	1 94	2 70	1 15

The following analyses by C. He is a h (Analysis, 142) show the range of variation of certain of the constituents of commercial rocated occoa-beans. The difference in the proportions of husk is due to the great variation in the thickness of the shells of occoas from different sources.—

	Propor	Roasted Bean after Removal of Husk									
Kind of Cocoa	Husk				ge	rates,	Ash				
	Par	Mohiture	Fat	Nitrogen	= Protesds	Carbohydrate: &c	Total	Sol 11a Water	в, Б.		
Carnones, .	18 8	4 32	48 4	1 76	11 14	92 10	3 96	2 15	154		
Trinidad (inferior),	15 5	8 84	49 4	1 78	11 14	82 82	2 80	0.98	0.88		
Surinam, .	15 5	3 78	54.4	176	11 14	28 35	2 85	0 80	1 23		
Gunyaquil, .	115	4 14	49 8	2 00	13 08	80 47	B 50	175	1 87		
Grenada,	14 6	8 90	45 6	1 98	12 40	35 76	240	6 80	1 85		
Bahra,	9.6	4 48	58 8	1 17	7 40	25 36	2 60	0.80	1.26		
Cuba, .	12 6	3 72	45 3	1.87	8 07	39 41	2.00	0 95	1 18		
Para,	85	3 96	54 0	2 00	12 66	26 34	8 65	1 40	1 00		

J. Bell (Analysis and Adults ation of Fools) gives the following particulars respecting the composition and the ash of cocca-nibs and busk —

		Per	100 Par Coons	ts of	Per 100 Parts of Ash						
	Kind of Coron	Mosture	Alkaloid.	Ash on Dry Substance	Soluble m Water	Insol in Acad.	P20s	CO2	K ₂ O.	¥e0	
į	Guayaquil niba,	5 08	0 54	8 63	56 20	none	40 39	0 60	28 85	0 21	
į	Surinam mile, .	4 55	9 80	2 20	43 46	none	27 78	3 81	28 00	0 88	
į	Grenada nibs, .	571	0 01	2 82	48 58	none	30 20	2 92	27 64	0 15	
į	Finest Trimdad nibs,	1 47	0 84	2 75	1G 55	Bone	89 20	4 10	20 80	9 11	
i	" " husks,	10 19	1 85	S 63	51 92	6 91	17 17	10 80	87 89	0.63	

In these analyses the figures for alkaloid are probably considerably below the truth

The ask of cacca is distinguished by the small proportion of chlorides, carbonates, and sodium compounds contained in it, and by the great prepondorance (3 or 5 1) of magnesia over line

In Bell's analyses of occos-ash, no mention is made of the presence of copper Duc laux proved this medal to be constantly present in cocoa. Galippic confirmed this, and found proportions varying from 00112 to 00939 gammes, pot longing manne of cocoa. The greater part of the copier existed in this makes, and in infrore kinds of chocolate containing cocoa-habit in large proportion copper was occasionally present to the extent of 0126 gazame per kindgramme.

The most important and characteristic constituents of cosca to the alterior them are a small proportion of cassens resources are small proportion of cassens resources present in addition. The necoviced proportions of theorems are very variable and generally unbrustwothy. This method of determination has already been described (nege 490). P. Troganowski (Archive der Pharm, [3] x 32, Joss Chem cosc, xxxxx. 853 from from 1 2 to 4 6 per cont of theoremsume in cocca, and concluded, from the results of a large number of experiments, that the proportion of alkaloid does not always bear a relation to the quality and value of the cocca. This is probable, but the difficulty attending the accurate determination of the colorisms and deduction of the kind of vey doubtful value

The fat of cocoa (Oleum Theobromatis, BP), sometimes called "cocoa butter," consists chiefly of stearn, and is fully

described on page 568. The proportion of fat present in coconnibs, free from husk, varies only a few units on each side of 50 per cent, and hence is valueless for the discrimination of samples from different sources

The taste and aroma of cocoa are chiefly due to a volatile substance, probably an essential oil, which appears to be developed by reasting, in the same manner as the caffeel of coffee (page 532) The tanish of cocoa also contributes to the flavour

The cocca-red probably does not pre-exist in cocca, but is a product of the oxidation of the tannin. If cocoa, from which the fat has been previously removed (by petroleum spirit), be exhausted with alcohol, and the solution treated with acetate of lead, a precipitate is produced, which, when suspended in water and decomposed by sulphuretted hydrogen, yields a clear and colourless filtrate, but on evaporating this hound, it acquires a bright red colour, and on taking up the residue with water, cocoa-red remains msoluble Cocoa-red gives various coloured precipitates with metallic salts, the tints depending on the extent to which oxidation has occurred, and, apparently, on the variety of cocoa employed. P. Troganowski (Aschw. der Pharm, [3], x 32, Jour Chem Soc, xxxn 363) has described various coloni-reactions yielded by the aqueous or alcoholic solutions of cocoa from various sources, but the value of the indications obtained is very questionable

The qum of cocca closely resembles gum-arabic in appearance, and yields mucic acid on oxidation with nitric acid. It differs from gum-arabic in being strongly dective-rotatory

The starth of cocon is present in only moderate proportion, and the amounts recorded by some observers are probably in excess of the truth. The granules are small, round, and exhibit a central hium. Under the microscope they are readily distinguished from the granules of added starbles.

Nitrogenous constituents of cocae G W Wig p or (1878) showed that of the mixegen of cocae only a pation varying from 39 to 73 per cent causted in a cognidable form (Analyst, v 8). The total introgen, as determined by combination with sode-line, ranged from 0.70 to 2.98 per cent, and that causting as coagniable alluminous from 0.33 to 2.35 per cent. According to Wigner, of the mixegen in a non congrubable form, part cents as theoremme and a furthen portion as nitrates. Wigner agued from this that the value of cocae as tool had been over-estimated

Weigmann similarly found only 42 per cent of the nitrogenous substances in cocoa to be digestable, and Stutzer states that, in spite of apparently favourable conditions, due to the physical condition of communical cocoa, a large proportion of the nitrogenous constituents remains entirely indigestable. Statzer classifies the nitrogenised compounds of cooca as follows:—I Non-protends, substances soluble in neutral aqueous solution in presence of entire hydroxide (theobromine, ammonis, amido-conspounds). 2 Digestable albumin, insoluble in neutral aqueous solutions in presence of entire hydroxide, but soluble when treated accessively with and gastic pures and lightine paneross extract

3 Insoluble and indigestable nitrogenous substances

The following are the results of the analysis of four cocopowders assumed by Stutz sor (Zeithot F angese. Chem. 1891), page 368) for the purpose of determining the effect of the process of manufacture on the chemical constituents A was composed of 40 per cent Aribs, 40 of Machais, and 20 of Bahna cocos, and was manufactured by Witskop & Co without the use of chemicals. B is a sample of a well-known cocon manufactured in Holland with the slightness of poiss. Consideration of the control of the significance of the control of the slightness of the

	A	В1	c	D,
	Per ernt	Per cent	Per cent	Per cent
Water,	4 80	8 83	6.58	541
Filtre,	8 30) 38 02	87 18	29 90	36 00
Total nitrogenous substances,1	30 84	19 88	20 03	19 25
Fut,	27 88	30 51	27 84	88 65
Ash,2	506	8.80	6 18	5 13
	100 00	100 00	100 00	100 00
1 Containing total nitrogen, Composed of —	3 06	8 80	8 06	8.57
Theobromine,	1.02	173	1.98	180
Ammonia,	0.06	0.03	0 40	0.33
Amido-compounds,	1 43	1:25	9 31	1 31
Digestible albumin,	10 25	7 63	10 50	7 81
Indigestable nitrogenous substances, .	7:18	9 19	7 68	8 00
Containing aftrogen,	1 16	1 47	1-23	1 28
Proportion of total nitrogen indi gestible,	81.2	44.5	81.2	85 8
2 Containing -Total PaOs	1.85	2 62	2 14	2 05
P_2O_8 soluble in water, .	1 43	0.50	0 74	0 77
Ratio of total P2O5 to soluble,	100 77	100 19	100 . 31	100 37
Ash soluble in water, .	8 70	4 70	2 82	2 78
Ratio of total ash to soluble,	100 74	100 67	100 64	100 49

An analysis of the ash of Van Houten's eccon by Konig (in 1880) showed:—Total ash, 7.84, K₂O, 8.52, CaO, 0.27, MgO, 0.81; P₂O₂, 1.84

COMMERCIAL COCOA AND CHOCOLATE

In its simplest form, commercial cocoa consists of the roasted and hunked seeds ("mine") ground to a past or semi-fund, and run into the form of cakes Flake coor is sometimes made by passing the decortacted seeds through a particular kind of rollers, but it is mostly made from the small particles containing much shell and zerm, semanted by the serves.

The term "cocc" is sometimes misapphed to mixtures of real cocca with sugar, &c. The practice is highly objectionable and has led to much confusion. It is better to describe all such cocca mixtures as chocolars, reserving the name cocca for the unmixed article

All good coose preparatons should be made from the cotylectors only, though the hunks enter nut the composition of many of the inferior kinds of coose and chocolate. In Germany, under the name of "coose-tes," and in Rieland as "muserables," conlusts are an independent article of commerce, the infrason of which in bothing water is drunk after the manner of tea

The large proportion of fat in occos (averaging 50 per cent) renders it impossible to manufacture a permanent preparation in the form of powdes, without either removing a portion of the fat or dulting the material with some non-fatty matter, such as sign, starch, or farma. Hence, there are two distinct types of "cocos" known in commerce, namely ".

- Preparations commonly called "cocoa-essence," or "cocoa-extract," consisting of ground cocoa-nibs, from which a part of the fat has been removed by heat and pressure
- 2. Preparations to which sugar and, geneally, some starchly material larve been added The sugar is usually sucrose (cane or best sugar), but reducing sugars are sometimes present in notable quantity Of the pure starches, arrowroot and rose starch are used in the better preparations, while wheel and potate-starches and wheat-flour are also met with. Moeller also mentions scorn and rys flours, ground earth-nuts, and malt, to which M a of adds almond-asks and sawdust. Any cheap vegetable material, capable of being reduced to fine powder, is liable to be used by unscrupilous

per cent Belohubeck (m. 1888) found —Total ash, 788, and for 100 of total ash, X₂O, 52'39, CaO, 1'56, MgO, 1045; P₂O₂, 24 91, OO₂, 3 45 per cent 'In large cooon manufactornes the busks are sented by sioves mito several

ares The largest are employed for unfasson, and the finest, containing a considerable admixtme of the kernisk, are ground up with super and exceeded to produce a low grade of checolate. The intermediate was are not readily applicable for eather of the above purposes, and hence effect a lower price than the consent and finest limits. They are employed for extile food, and at Hambug are pressed for the activation of exceeded reading the contraction of exceeded the contraction

cocoa manufacturess, but the better class of preparations which have acquired a reputation in the United Kingdom are free from any suspection of such admixtures.

A considerable addition of cacao-butter is made to some kinds of chocolate.

The flavous ray agents added to chocolate are most frequently vanilla and cumanon. Artificial vanilla, unturing, gloves, moéce, are also used. In addition to the mechanical difficulty it manpulsing undinisted econe containing all its natural fast, it stated, with some probability, that the excessive proportion of the tanders the occos difficult of digestion. Hence the removal of a portion of the fast, and consequent concentration of the non-fatily constitutions of the coor, any parts to be distinctly advantageous

A further iteatment of the concentrated corea is practised by some manufactures of cooce-seemoe, sepsently by Dutch firms. This tractment consists in the addition to the cocea of an alkali, which may be either ammonia or a fixed latch or alkaline cobonate, whereby the fat becomes emulsified and any free fatty acids esponified. Hence, on subsequently teating the cocea with hot water there is less teadency to the separation of only globules. The effect on the composition of the cocea is shown in the results of Statize on page 560, from which it appears that the fact of the treatment can be resulty detected. In the case of a well-known brand of cocea, poissaum carbonate is used. In another case, the coceabeans are seaked in water containing from 2 to 4 per cent of their weight of cousts potsaol or code.

The following figures were obtained by the analysis, in the author's laboratory, of a specimen of the best eccoa-mbs and two of the leading brands of cocan-essence or soluble cocas, to which no starch or sugar had been added.—

	Coton	Sample A	Bample D
ASEmasteble in water, Soluble in water, OOLD HOY Containing —Ash, Organic extract,	Per cont 2 63 1 71 0 82 0 32 9 72 0 60 0 63 16 84 3 34 13 56	Per cent 1 93 3 50 1 13 0 40 11 01 0 71 0 70 20 36 4 03 15 43	Por cent. 8 25 2 00 6 10 5 03 18 00 2 02 0 34 27 16 7 86 10 31

^{1 &}quot;Chocolate creams" consist of a core or keinel of pure sugar, enveloped in a mixture of ground cocoa, cacac-butten, sugar, and flavouring materials.

The cursons property possessed by the cold-water extract of being at once alkaline to methyl-range and and to phenolphthaleum indicates the presence of a soluble sait of some weak organic acid, together with a small proportion of free organe send. The treatment with alkali which sample B had secoved appears to have notably incressed the proportion of matter actually soluble in water

The musues of the term "soluble" by cocca manufacturers is notorious, the real object scopit, and to some extent statused, being the formation of an emulsion which is readily muschle with hot water. This deader-durm is the more important owing to the difficult digestibility of some of the nitrogenous constituents of cooks (see page 559)

The following results, among many others, were obtained by E E we'll (Bulletin No 13, U S Department of Aguiculture) by the analysis of well-known brands of commercial cocca and its pre-narations.—

				ā		A	sit	
Description of Sample	Munk 1	Fat.	· Fibro	Cane-sugar	Reducing	Total	Acad equiva- lent 2	Added Struch
Fry's Cooos Extract	1	30 95	8 89	Г		4 21	58	Nono
Schweitzer's Coccatina,	1	81 13	3 70			6.33	94	None.
Van Houten's Cocos,	1	29 61	4 38	١.		8 64	16 05	None.
Blooker's Dutch Coccs,	0	\$1.48	8 76	ļ		6 96	96	None,
Rowntrae's Extract }	2	27 58	6 62		-	8 48	16 6	None
Rowntree's Powdered Chocolete,	2	25 84	1 38	51	none	1 08	2 25	Vory small amount of airowroat
Epps' Propared Cocca,	ŀ	25 94	151	26	none	8 15	26	Much arrowroot
Fry's Dismond Sweet }	2	18 00	81	55	some	1 16	145	Much wheat starch with some arrow-
London Cooss (nn- known maker),	8	11.13	2 13	82	some	282	89	Very largely diluted with amounted
Chocolat-Memer,	0	21 81	1 16	58	none	1 40	2 05	None

In the column handed "mink," 0 signifies that no pharacteristic bunk kause could be found under the microscope, a sugnifies that the laws had probably been madity temovarly. 2 signifies that the laws had probably been madity temovarly, as 2 signifies that the laws had probably been partly removed, and 2 that the laws have probably all propent. But is most to characters with support to the laws for commercial consens are not in all cases home out by the examination of other samples of the same proparation, and must be received with constitution.

² The figures in the column headed "acid equivalent" represent the number of c c of decharmal acid required to noutralise the ash from 2 grammes of the sample It is a rough measure of the fixed slikeli used in the manufacture.

Owing to a considerable proportion of the instant fat having frequently been removed, the proportion of tool coop in a mixture cannot be assumed to be approximately double the percentage of fat. A batter idea of the proportion of the adultions is obtained by stating the fat and non-fatly constituents separately. This plan is adopted by J. B. el I, and is shown in the following analyses by him, representing the composition of certain commercial preresponding of conosition.

Description	Moisture	Fat	Added Sugar	Added Staroh	Non Inthy Couga (by differ- ence)	Nittogen
Finest Trinidad nibs,	2 60	51.77	none	none	45'03	2 05
Cooos Extract, .	5 78	29 50	none	none	64 74	Not determined.
Flake Cooos,	5 49	28 24	none	none	66 27	8 08
Coxontina,	8 62	23 98	none	none	72 50	4 07
Chocolatine,	4 40	29 60	none	none	68 00	6 36
Chocolat de Santé, .	144	22 08	61 21	2 00	18 27	Not determined.
Prepared Cocon,	4 95	24 94	28 08	19 19	27 89	2 24
Iceland Moss Cooon,	5 47	16 86	29 23	24 70	23 74	1:88
Rock Cocon,	2.08	22 70	32 20	17 58	24 90	Not dotermined.

According to evidence given in the case of G i b s on v L o a p.e.r, "Speciocoan" contains 40 per cont of cocoa, 16 of starch (Westin Indian arrowroot), and 44 per cent of supar "Ghamilated cocoa" is cheefly a mixture of cocoa-mile, sugar, and arrowroot, while in "Maravilla cooa" the sirrowroot is replaced by ago. B er n in a r dt states that be has met with chocolates commenting of cocoa-romnants, fat, sugar, spaces and colouring matter, and containing no true occoa whetever. The cocoa-butter is said to be liable to be replaced by cheaper fats, and vanilla and vanillin by Peruvian or Tolin balsam, storms, or your business.

ANALYSIS OF COMMERCIAL COOOL AND CHOCOLATE.

The complete analyse of coosa is rawly required. A careful microscopic examination will undeate the presence, and in many cases the nature, of most foreign additions, and prove the presence of hust-structure. The various starches may also be identified by the nucerospore. The proportion of fait affords further information, and the percentages of sugar and starch complete what is usually required, unless it is desired to ascertain the nature and amount of the alkah added. The following scheme of nautres will allow of the above information here obtained on analyses will allow of the above information here obtained.

Ignite 5 grammes of the sample, weigh the ash and treat with boiling water Wash, dry, ignite, and weigh the insoluble portion Titrate the filtrate with decinormal acid to determine the alkalinity, which will be excessive where the cocoa has been prepared with a fixed alkali The addition to cocoa of ferruginous pigments, such as rouge, othre, and venetian-red, was formerly practised, and the author was recently consulted as to the probable legal consequences of their use. He has also examined a preparation consisting essentially of oxide of non, which has recently been offered to cocoa-manufacturers Where the proportion of the diluents is large, the importance of deepening the colour of the mixture is considerable. The addition of ferruginous matters would be readily detected by the excessive proportion of the ash, which in the case of genuine cocoa is white, and very rarely in excess of 4 per cent (in the absence of husk and added alkalies, and when the fat has not been removed) The proportion of oxide of iron in cocoa is very trifling, ranging from 010 to 038 per cent of the ash, while even in the husk it only amounts to 0.63 per cent of the ash

Dry 5 grammes of the sample in the water-oven at 100°C and note to loss of weight, which represents mosteries. Boil the dract substance, reduced to powder if necessary and preferably mixed with a known weight of dry sand, with reducibilet, petroleum spirit Pour off the solution, and report the treatment till the fait is circle; removed. Weak the isselined, grif in the both and, roweigh. The loss represents fat, with a near approach to account a direct distance of the result of the result

The residue left after the extraction of the fat is exhausted with hot spirit of 0.850 specific gravity, which dissolves sugar, tartaric theobromine, &c The hot solution is treated filtered from the precipitate of lead tartrats,

tannate, stearate, &c. From the concentrated filtrate the Harbromine can be extracted by agutation with warm chloroform, but where the determination is not required this stage of the pieces may be omitted. The aqueous liquid is freed from traces of chloroform by boling or egitation with periodium spirit, and after removal

Cocon which has been tested with an alkali contains a notable quantity of cose, which is not dissolved by the petroleum edie. It is best extended by tracting the reastine with alcohol containing a few drops of hydrochical containing a few drops of hydrochical way containing the alcoholo containing a few drops of hydrochical way contained the containing the solution is contained and the residual luquid with water and other. On separating and evaporating the othered layer, the fatty ands of this cosp will be left.

of the excess of lead by sodurm phosphate is fit for determination of the sugar. This may be effected by inversion and treatment with Fehlung's solution, or by means of the polarimeter. The difference in the amount of sugar found before and after invorsion represents the came-sugar added. The alcoholic extinct of genuince cook, after treatment with lead acetate, does not sensibly reduce Fehling's solution, so that any preceptate prelded before investion represents glazzes, introduced as such or present in the came-sugar addict.

The resulue left after treatment with alcohol contains gum, starch, collulose, fibre, alloumnoud matters, &c. After weighing, an aliquely part may, if desared, be used for the determination of the contained introgen by Kjoklahl's process or combustion with sold-time, and the amount found calculated to allounized by multiplying by 6.25. The results may also be advantageously examined under the microscope at this stage, since by the removal of the oil, sugar, and colouing matters the starch and woody structure are seen to great advantage. On the presence of alsence of foreign starch will usually depend the necessity of performing the subsections operations for its quantitative determination.

For the determination of starch, an aliquot part of the residue from the alcohol treatment 2 should be heated, under a pressure of

1 A determination of the amount of sugar added to coops can be readily effected to within 2 per cent of the truth, but a strictly occurate estimation is not required, and would be very difficult. The sugar can be determined in the aqueous instead of the alcoholic extract of the cocea, but in that case the solution contains the natural gum, which has a doxino-rotatory power equivalent to 0 3 to 2 0 per cent of cano-sugar in the sample, and a large volume of cold water must be used for the overaction. E E Ewell (Eulleten No. 13, U.S. Department of Agriculture) recommends the following method for the polarimetric determination of angar in the aqueous extract of cocon -18 024 grammes weight of the material is triturated in a small mostar with alcohol until a smooth paste is obtained This is transferred to a 500 c c. flask. diluted with about 400 c.c. of water, and the liquid shaken occasionally for three or four hours, when 10 c o, of a saturated solution of mential lead noctate should be added and the volume brought to 500 c c. After standing for an hour with occasional agrication, the solution is filtered and policised in a 4 decimetro tube (twice the usual length) If the instrument be one intended for use with 26 048 grammes of sugar, the percentage of cane sugar in the sample will be found by the following formula, in which R is the reading in sugar-units ---

$$\frac{R}{100}$$
 $\left[600 - (18\ 024) - \frac{5R \times 13\ 024}{100} \right]$ - per cont of sucross,

² The residue is preferably first treated with cold water, to disselve gummy matters, but except in cases where great securacy is required this part of the process may be omitted.

1 atmosphere, for one hour with 50 c. of water and 1 c. or of truming hybrothore acul. This teatiment effects the complete conversion of the starch into multiose and dextrin, and the further change of these to dextrose, without appreciably affecting the callulose. The solution is filtered from the mediulose matter, fibre (sand), &c. and the dextrose determined in the neutralised fibre by Fehling's solution. Ten pairs of dextrose found represent 9 of starch in the samele

The mixed colludes, flore, and sand, left after the conversion of the statch by hydrochloric acid, should be treated with a solution of 2 per cent caustic soda to temore introgenous matters, washed successively with very dilute hydrochloric acid, alcohol and ether, dread and washed 2

An alternative method of estimating storch consists in treating the fat-free coors with cold water, to remove all sugar, gum, for. The liquid is filtered and the readue washed with decinormal coasies ods of 4 grammes NadHo per litre) to remove albousimeds. The readue is rinsed off the filter with warm water, the liquid heated to boiling while constantly stirred, so as to golatimes the starch, and the product treated with a known measure of recently-prepared and filtered cold aqueous infusion of malt, of which the specific gravity has been previously ascertained. The mixture is kept at a temperature of 60° to 63°, with occasional stirring, until a drop taken out with a glass rod and sided to a drop of ditte todine solution on a porcelain plate shows no blue or brown coloration. The solution is then filtered, made up to a definite volume,

A supple and convenient apparatus for effecting the convenient constants as a soft-water bottle field with an indicatible stopper, through when between a long glass take best twice at right angles and amoremed to a depth of 30 makes in menery contained in a long vertical glass in the or pace of all of arrow (roa) gas-rape. The stoppes should be carefully secured by wire. The soft-well posterior of the stoppes should be carefully secured by wire. The soft-well posterior that the present in oil, or in a botting saturated aqueous solution of sodium initiate. This last input has a temperature of 121° O, corresponding to one additional atmosphere of piessure, so that no regulation is required, and if preferred the exti-tuto may be discussed with the solution of the contraction of the co

* For the direct determination of the oracle flow, 2 gammes of the ample occose should be freed from fit and botale for half an hour under a sight conclusion with 500 c.c. of water and 2½ c.a. of sulphure sold. The buguet is effect through home and the reaches thereogally washed with hot water and then botical with 200 c.o. of 1½ for cont. counts sold. The reaches soffices of (washed in secseons with hot worts, alcoloi, and atther, direct 2.110, and free from bank it will amount to 2 or 8 per cent. only, but will exceed this limit in prospection to the amount of Aust Process.

and its specific gravity accurately ascertained. From the excess of the density over vater a subtaneted the density due to the infusion of malt used, allowance being made for the increased volume of the liquid, when the difference represents the density due to the stanch dissolved, and thus number divided by 4 098 (= 3 95, the density-coefficient of a solution of mixed maltices and dextrim, multiplied by 1 037, the yield of these from 1 part of starch) gives the number of grammes of starch in each 100 c.c. of the solution.

The total sutroges of cocos can be determined on 2 to 3 grammes by Kajelaha's method, or by combuston with soci-time, The assumption that the proportion of albuminous can be found by multiplying the nitrogen by 6 25 leads to an estimate greatly in-excess of the truth. The theobtomine of cocos contains 31:1 per cent of introgen, or nearly twice is much as albumin. Hence to obtain an estimate of proteads from the introgen of the sample, the proportion of that element corresponding to the theobtomine present must first be deduced. But as the determination of theobtomine is somewhat troublesome, it is preferable to operation a cocos-readened which has been already exhibited with the protein spirit, alcohol, and ample alcohol or chloroform, so as to eliminate with certainty the whole of the theobtomine.

CAGAO-BUTTER (Oleum Theobromatis) is the fat contained in coccabeans, and must not be confused with cocoa-nut oil from Cocos numbers

Cono-butter is expressed from econo in the process of manufacture, and by far the larger ganantity used in the United Kingdom is the produce of one firm. It is used in pharmacy, for the production of some kinds of chocolate, and in the manufacture of high-class soap. Cono-butter is hable to adulteration with or substitution by other fats, and it is said that the ocano-butter is sometimes very completely expressed from coons and replaced by tallow, coco-articl, or other comparatively cheap fat

A careful observation of the physical and chemical characters of

density of the solution less that of water (1023-1000-23), leaves 20 as the occess-density caused by the solution of the starch of the sample, and thus figure divided by 4 998 gives 4 9 grammes per 100 a.o or in the 20 grammes taken, or 24 5 per cent of starch in the sample

cacao-butter will allow of the detection of other fats, if present in any considerable proportion

Pure eases-butter is a yellowsh fat, gmdually becoming paler on keeping. I At the ordinary temperature it may be broken into fragments, but softens in the hand and melts in the mouth. Ciccabutter has an agreeable odour, tastes like choolate, and does not readily become raned. It basedwas m 20 parts of het alcohol, separating almost completely on cooling, and is also soluble in ether, section exists when the superior of the cooling and is also soluble in ether, section when the superior was superior when the superior was a superior when the superior was a superior was a superior was a superior when the superior was a superior was

Cacac-butter owes its value for the production of pessance and suppositories to the fact of its having a multine-point slightly below the temperature of the human body (98° F = 36·6° C). According to most observers, it fases between 30° and 33° C (travely at 26°) to a transparent yellowah liquid, which congoals again at 30°-21°, the temperature range to about 27° C. According to the British Pharmacopiesa, the melling-point of coacac-butter range between 30° and 35° C (68°-30° F) §

¹ It is to be regnetted that the yellowish tint of cacao-butter is not more generally incognised as a natural characteristic. It is probable that the quality of cacao-butter is necessarily affected for the worse by any process of decolorsistic.

¹ R Beaseman n (Seat Anal Chem, xxiv, 283; Jose Soc Chem Inc., to 585) has observed the melting-point of casso-butter and the fatty scale resulting from its suppositionation, and finds the figures for the latter sensitive and the state of the present of the present of the present of the state of the winds part of a piece of quill-tubing diam's not to a capillary form and closed at one one. The substance is allowed to soldity completely, and the tube is then statemed became it is greated by the state of the state of the state of the state of the present of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of in a cylent of the state of the point of the state of the point of the state of the point of the state of the state of the point of the state of the state

	Fat	Patty Acids						
Source of Cacao buttes	Initial Melting point	Initial Melting point	Concluding Multing point	Percentage of Insoluble				
Maracatho beans, .	25-26° C	48-40° C	61-62° C	94*89				
Caraccas beans,	27-28	48-40	51-62	95 31				
Trinidad beaus, .	28-27	49-50	52-53	95 65				
Portoplata beans, .	28-29	49-50	52-58	96-48				
Machala Guaysquil	28-29	49-50	52-53	95 24				

T M Clague has recently pointed out (Phaim Join [3], xxin 247) that the melting-point of commercial cacac-butter extends over a considerably greater range than the above, and is materially affected by the temperature to which it has been exposed. Thus, the melting-point of ten trieds samples ranged from 78-91° F A sample expressed by heat direct from coose-mbs mulcled at 91°. A sumple oxpressed by heat direct from coose-mbs mulcled at 91° and the time of the melting-point of 96°, while the care-butter extracted by host and resulting-point of 96°, while the care-butter extracted by host and pressure by the same firm molled at 75° F, thus showing that a certain amount of fractionation occurs in the ordinary process of extraction by pressure.

T. M. Glag us further observed the following suggested to alterations of multime-point when concolution was heated to various temperatures. Nos 1 and 2 were ordinary trade samples, and hence had been already heated in the process of manufacture. No 3 was extended by either from uurousted occosaribs, and hence accessive heating had been entirely avoided.

									Melting point, "F		
									No. 1	No 2	No 3
Original,									76	88	84
After being	heated t	io 105° P						-	75 5	80	86
**		120*,						1	81	85	92
**		150°,							83	88	92
15	**	180°,						-	80	80	85

The melting-point of No. I sample was raised to 86° F, by keeping it at a temperature just under 100° F for two hous. The determinations of melting-points were made on metallic mercury, substantially by method e described in Vol II page 23.1

Connobutive contains the givenness of steams, oles, and a hitible haure, palmite, and arachide acids. C. T. K. in great to obtained from connobutive an acid of the formula $C_{\rm eff} = 0.0$, which he named the obromic acid. P. Graf modulo 959 per cent. of givenol, and detected a hitle cholesterin and small quantities of formus, aceies, and buttern ends.

¹ T M Clague has also described as paraments showing that determinations of the melting-point of cacao-butter by the capillary tube method are very gravely affected by the diameter of the tube employed. The todine-absorption of a large number of samples of concolutior from different sources has been determined by F filsinger (Chem Zest, xiv 716), and found to range from \$3.4 to 37.5. The supoinfleation-equivalent ranges a few degrees on each side of 250, which liguin corresponds to 20.03 per cent. of potash (KHO) required for suportification. Filsinger found in potash required to lange from 19.2 to 20.2, and Weigin an inform 19.84 to 20.30. An admixture of paraffin wax would reduce the percentage of alkalt required for suponfleation.

The specific gravity of solid cacao-butter is variously stated. The author found the plummet-gravity at 98° C to be 0 8577. Any admixture of parafiin wax would reduce this figure, while coccanut oil would increase it

Foreign fats in eaco-butter tend to alter the foregoing characters, but observations of the melling-point and specific gravity do not usually furnish satisfactory means of detecting such admixtures. Tallow is said to be capable of detection by saturnting a cotton thread with the oil, allowing it to burn for a short time, and then blowing it only, when the odour of tallow becames precentible.

A better test for tallow and other adulterants of cacao-butter is to dissolve 2 grammes of the fat in 4 grammes (=55 cc) of ether at 17°-18° C,1 and then immerse the closely-corked testtube in ice-cold water Granules will separate from, or turbidity be produced with, pure cacao-butter, in not less than 3 and more frequently in from 5 to 8 minutes, sometimes delayed to 10 or 15 minutes, while if tallow or suct be present, a turbidity will appear at once, or within 24 minutes, according to the proportion of the adulterant, of which 5 per cent may thus be detected On exposing the solution to a temperature of 14° to 15°, it will gradually become clear again, or more rapidly at 20°, if the cacao-butter was pure, but not if it was adulterated With a sample containing 5 per cent of tallow, turbidity occurs in 8 minutes, and the solution does not become clear below 22°, while with 10 per cent of tallow, the turbidity occurs in 7 minutes, and the clearing-point is 25° C This test is due to Bjorkland (Zeit Anal Chem, in 233), and is adopted in the United States Pharmacopona Its value has been confirmed by other observers, of whom Lamhofer has pointed out that petroleum-ether may be employed with similar results, except that the cacao-butter separates rather more slowly than from other, the deposit being always granular, while other fats render the entire liquid cloudy. The solution of cacao-butter in two parts of ether will remain clear for an entire day if maintained

¹ A failure to obtain a clear solution points to the presence of paraffin wax.

1. 2.

at a temperature of 12° to 15° C. This modification of the test is prescribed by the German Pharmacopæia, and is due to Ramsperger, who states that aniline may be substituted for the ether. Filsinger (Zest. Anal. Chem., xix 247) has described the following modification of the ether-test -Two grammes of the fat should be melted in a graduated tube with 6 cc of a mixture of 4 volumes of ether (sp. gr. 0 725) and 2 volumes of alcohol (sp. gr. 0.810), shaken, and set aside. Pure caouo-butter gives a solution which remains clear

According to E. Dietrich, a very reliable test for the purity of cacao-butter consists in warming the sample with an equal weight of paraffin oil. A drop of the mixture is placed on a slip of glass, a thin cover applied, and the slide exposed for twelve hours to a temperature not exceeding 5° C. When then examined with polarised light, under a magnifying point of 20 diameters, the crystals of cacao-butter present the appearance of palm-leaves, showing a fine play of colours with selenite. An addition of 10 per cent of beef-tallow causes the fat to crystallise in tufts of needles, or circular groups of crystals, which exhibit a black cross; while if mutton-tallow be the adulterant, it is stated that no cross can be seen.

References to Photographs of Leaves. (See page 522)

PLATE I. PLATE II.

	14,	Camellia Thea. Tea.
Marattia Elegans	15.	Ribes Grossularia Gooseberry,
Bpilobium Angustrfolium. French	16	Rosa Canina Dog Rose,

Willow or Willow Heab. 17, Coffee Arabica, Coffee 4. Salaz Alba. Willow How 18. Crategus Oxyacantha Rez Paraquanensis Paraouav thoin. Tes or Brazilian Holly 19. Fragaria Vesoa. Strawberry.

Populus Nigra Poplar 20. Pyrus Malus. Apple. 7. Sambuous Nugra. Eldor. 21. Quereus Robur. Oak 8. Ulmus Campesty 1s. Elm. 22. Ribes Nigrum Black Current.

9. Betula Alba Buch 23. Fraziniis Excelsior. Ash 10. Prunus Spinosa. Sloe or Black-24 Fagus Sulvatica, Beech. thorn.

Rubus Fructicosus. Black-11. Prunus Cerasus, Cherry pen A

12. Rubus Idaus Raspberry. 26. Prunus Communis Plum. 13. Camellia Sasanqua.



INDEX.

AGREAMING 2 Acid, matropic, 246, 286 Acetanilide, 63, 67, 68 ---- 180cocate, 287 --- detection of, 69, 83, 84 --- isonicotimo, 111 Aceto-amidophenol, 68 - kinto, 445 Aceto-anisidine, 68, 85 - leucotronic, 263 Acatophenone, 68 - Intidinic, 112 Acet phenethidine, 68, 81 - lycootonic, 225 Acet-phenylhydramae, 28, 68 - meconic, 386 Aoid, acomitto, 207 detection of, 388, 858 ---- amalic, 480 - metatungstio, as a reagent, 137 - amidobenzene-sulphonic, 49 - methylparabanic, 494 - amidonaphthol-sulphonic, 94 --- nicotinic, 111 - amino-sulphonie, 49 - mitric, as a reagent, 146 --- atrovio, 245 - opianic, 203, 298 --- berberonio, 112, 468 - phosphoantimonic, as a reagent ---- bohere, 501 --- cafferc, 529 ---- phosphomolybdic, 136 - caffeidine-carboxylic, 479 - phosphotungstic, 136 - caffeiannio, 529 - picolimo, 111 - cholidonic, 166 - picrio, as a reagent, 134 ---- chlorogenic, 529 - pyridine-carboxylio, 110, 112 --- ohrysatropic, 262 - pyromeconic, 837 ---- omchomeronic, 112, 165 ---- quinio, 445 - cinchofulvio, 445 - quinolinic, 112 _ -- quinovic, 444 - silicotungstic, as a reagent, 187 --- cocatanuic, 291 - strychnic, 384 - strychnine-monosulphome, 363 --- columbio, 472 - sulphanilic, 49 --- comente, 887 - cotarnic, 299 - sulphomolybdic, as a reagent, 147 ---- dimethyl-parabanic, 481 - sulphoselenic, as a reagent, 145 dipicolinic, 112 - sulphovanadic, as a reagent, 148 ----hemipinic, 298, 463 - sulphuric, as a loagent, 145 - hydrochloric, as a reagent, 145 --- tannic, as a reagent, 135 . - hydroquinine-sulphonie, 425 ---- tannic, in tea, 491, 515 - hydrastinic, 470 - taopic, 245 ---- 1gasunic, 884 --- tumethyl-thiocai bamic, 16

574 INDEX.

074 INI	INDEX.	
Acel, urio, colour-reaction of, 473 Acalysture, 224 Acounte, 202, 204, 214, 222, 285 Acounts, 202, 204, 214, 222, 285 Acounts, seasy of, 228 — official preparations of, 199 — passoning by, 298 — toxicological detection of, 240 — various species of, 199, 201 Acountine, 207 — smail-plossin, 201, 215, 218 — shiptine, 206, 233 — observation of, 208 — constitution of, 208 — constitution of, 208 — constitution of, 208 — determination of, 201, 203 — determination of, 201, 203 — determination of, 201, 203 — determination of, 201, 202 — bularyour of, 201, 202 — bularyour of, 201 — bularyour of, 201 — of bularyour of, 201 — of bularyour of, 201 — of calendine, 203 — of bularyour of, 201 — of calendine, 203 — of calendine, 203 — of calendine, 203 — of calendine, 203 — of calendine, 201 — of	Albalends of Remijia barks, 384 — outd-tion colour-seachous for, 149, 381, 988, 469, 460 — phy-sologocal tests for, 149 — purification of, 162 — seachous of, with Commpelités creeges, 144 — Diagendos if seages, 147 — Diagendos if seages, 148 — ficture thioule, 183, 304, 318 — Fulción reages, 147 — Jugordos reages, 147 — Jugordos reages, 147 — Jugordos reages, 147 — Mayar vinegos, 138, 138 — Mayar vinegos, 138, 138 — Marar vinegos, 138, 138 — Marar vinegos, 138, 138 — Marar vinegos, 138, 139 — Marar vinegos, 138, 139 — Marar vinegos, 138 — Subalber vinegos, 138 — Subalber vinegos, 138 — Subalber vinegos, 138 — Woundry's reages, 137 — rine chlorole, 141 — stry hime, 369 Altha-maphthylamino, 60 — phylithers, 94 — naphthole-sulphone ands, 40 — naphthole-sulphone ands, 50 — peans of the phylithenes, 69 — phenole, 80 — thosphane, 68 Amune, clevefecter of the destination of research of the destination, 60 — phenole, 80 — thosphane, 68 Amune, clevefecter of research of the destination of the phylithenes, 60 — hopehole, 60 — destination of, 7	
general preceptants of, 184, 158 of hemlock, 171		
of henbane, 250, 267	- separation of, 4	
of hydrasta, 461, 467	Ammol, 15	
isolation of, 151	Ammonium bases, 18	
of hpine, 178	Analgean, 32	
mydriatic, 244	Anhydro-acoustmo, 205, 213	
of nux vonner, 384		
of opum, 293	ecgenine, 251	

575

1-1-1 t t to	
Anhydro-tropines, 251	Bromscotamhdo, 68, 71
Antifehiin, 68	Bruome, 381
Antithermin, 31, 68	constitution of, 168, 381
Antipyrine, 32	dimitro-, 382
chloral-, 88	teachons of, 382
nitroso;, 84	soperation of, from strychnino,
Antiseptin, 68, 71	366
Apo-aconitine, 213	Butylamine, 14
Apo atropine, 251	
Apo-bases, see Anhydro-bases	CAPPRIDINE, 479
Apocodeme, 824	carboxylio acid, 479
Apomorphine, 319	Caffeins, 474
Arginine, 167, 178	assay of tea for, 490
Arieme, 393, 436	- constitution of, 167, 478
Atisine, 226	- determination of, 484
Atropamine, 244, 251	natual occurence of, 474
Atropino, and its allies, 243	presence of, tn cocca, 495
anhydro-, 251	proportion of, in tea, 492
constitution of, 165, 244	n coffee, 528, 552
products of saponification of, 244	reactions of, 480
reactions of, 254	salts of, 482
toxicological detection of, 261	solubilities of, 477
Azobenzeno, 68	Calumba 100t, 471
Azo dyes, 42	Camphor, compound functure of, 858
Azormide, 24	Canadine, 470
·	Carbazol, 118
Bases from tea, 39	Cevadino, constitution of, 183, 166
Belladonns, alkaloids of, 263	Cevine, 133, 166
assay of, 284	Chanamidino, 398, 436
composition of, 282	Chanamine, 393, 486
extract of, 289	Chelidonine, 295
Belladonnine, 244, 252	Cholerythine, 295
Benzanilide, 72	Chicay, composition of, 538, 544
	detection of, in coffee, 540
Benzdme, 88	determination of, 542, 545, 550
Benzoyl-aconino, 207	Chinoline, see Quinoline
	Chinovin, 448
	Chloral-antipyrine, 38
	Cholestrophane, 481
	Chocolate, 561
	Cholmo, 18, 193, 167
Benzylamme, 51	Cinchamidine, 433
Burbamine, 461, 466	Cincholenpone, 168
Beiberme, 461	Cinchona alkaloids, 391
salts of, 464	amorphous, 433
Berheris alkaloids, 461	general properties of, 394
Beta-naphthylamino, 90, 92	proportion of, in bark, 445

010 INI	DE.A.
Cinchons alkaloids, separation of, 453	Cocos, composition of, 556
list of, 892	essence of, 561
barke, 440	- husks, 556, 557, 561
assay of, 449	mtrogenous constituents of, 556
oomposition of, 442	Codamino, 294, 801, 804, 320
proportion of alkaloids in,	Codame, 294, 321
445	Coffee, 527
Cinchonamine, 892, 486, 438	- adulterations of ground, 538
Cinchons-red, 445	beans, 533
Cinchonicine, 485	- caramel m, 539
Cinchonidino, 892, 897, 428	composition of, 528
constitution of, 168	detection of chicory in, 540
- determination of, 410, 413, 480,	detection of starch in, 541
449, 459, 460	factitious, 535
homo-, 430	imitation, 535
hydro-, 432	
Cinchonine, 892, 397, 481	physiological action of, 532
constitution of, 168	roasting of, 530
decomposition-products of, 168	Colchicine, 166
determination of, 413, 459, 460	Collidines, 97, 109
hydro-, 392, 432	Columbin, 472
Omehotannin, 444	Concharanino, 898, 436
Cinobateniue, 168	Concusconine, 893
Cinchotane, 892, 432	Conhydrine, 171, 173
Cinnamyl cocame, 271, 285	Contcernes, 174
	Comme, 171
Coon, alkaloids of, 270	assay of hemlook for, 178
- amerphous bases of, 287	determination of, 176
leaves, 290	potsoning by, 175
- extraotion of alkaloids from,	Conum, 176
292	- alkaloids of, 171
Cocaine, 273	tinoture of, 177
- amerphous, 287	Conquinamine, 372, 427
- onnamyl-, 285	Conquimme, see Quinidino
commorcial, 278	Cotsimno, 299
constitution of, 166, 271	Ciyptopino, 294, 301, 304, 324
' '- 1 of, 282	Cumidinos, 59, 63
, °	Cuproine, 392, 397, 438
hemo-, 285	constitution of, 169, 398, 489
hydrochloride, 277	separation of, 413, 438
- saponification of, 272, 282	Outare, 887
Cocamina, 271, 272, 286	Curame, 871, 389
Coosthylme, 285	Caume, 390
Cocoa, adulterations of, 561	Ouscemidine, 393
analysis of, 564	Cuscamine, 893
commercial, 561	Cuscondine, 393
	Cueconine, 893

INDEX 577

Durana anti-n cor cor	P l		
DEUTEROPINE, 295, 324	Exalgin, 68, 71		
Diamide, 22	Extract of acouste, 229		
Diamidogen, 22	belladonna, 266, 269		
Diamido-henzenes, 86	emchona, 445		
Diamido-phenols, 83, 85	cocoa, 561		
Diantipyrine, 31	coffee, 553		
Diazo compounds, formation of, 7	hemlook, 77		
Diemehomeme, 893, 435	henbano, 269		
Diethylandine, 73, 79	nux vomica, 886		
Diethylene-diamine, 2, 106	opium, 350		
Diethyl-hydrazme, 26, 27	tea, 505		
Dimethylamine, 12	FLAVANILINE, 69		
isactions of, 10	Formyl-paraphenethidino, 85, 373		
Dimethylandine, 73, 74	Furtman, 113		
commercial, 76			
Dimethylnitroeamine, 7	GILPEMINE, colour-reactions of, 145,		
Dinaphthylene diamines, 93	146, 149, 487, 469		
Dinitiohenzenes, 63	Glancine, 298		
Diphenylamine, 79	Glaucopictine, 296		
Diphenylaniline, 79	Ghicosides, behaviour with immiscible		
Diphenylone-diamnies, 86	solvonta, 158, 159		
Dignimicine, 398, 435	colour reactions of, 146, 147, 148		
Dita baik, 436	Gnoscopine, 294, 801, 324		
Ditaine, 436	Cinotopino, 204, 001, 024		
	Henry one seems of 178		
Ditamine, 486	Hrmlock, assay of, 176		
Ditoinylene-diamine, 87	lesser , 175		
Diurotan, 497	poisoning by, 175		
7	- tmeture of, 177		
EASTON'S SYRUP, 376	water-, 175		
Echitamine, 436	Henbane, alkaloids of, 250, 267		
Echitemne, 486	assay of, 267		
Ecgonine, 283	extract of, 269		
auhyd10-, 284	Herepathite, 138, 402, 454		
benzoyl-, 282	Homatiopine, 258, 254		
constitution of, 166, 270, 284	Homoquinine, 439		
Ethyl diethyloxamate, 5	Hydracetin, 28, 68		
— oxalate, 5	Hydrastme, 461, 467		
Ethylamino, 14	colour-reactions of, 469		
reactions of, 10, 17	Hydrastume, 470		
Ethylamines, 17	Hydrazme, 22		
Ethylaniline, 73	ethyl-, 26		
Ethyl-hydrazme, 26	hydrate, 22		
	phenyl-, 27		
	salts, 23		
-thalline, 121	Hydrazines, 22		
Euphorm, 68, 72	- enbstituted, 25		
- 4 1	0.		

Hydrazebanzene, 89	Lobeline, 195
Hydrazones, 30	Loganetin, 385
Hydrazonium compounds, 25	Loganin, 385
Hydroscridine, 126	Lupanine, 179
Hydrocmehonidine, 392, 410, 480	Lupine alkaloida, 178
Hydrocinchonine, 392, 432	Lupindine, 179
Hydrocotarmne, 294, 301, 804, 309,	Lupinine, 167, 178
325	
Hydrohydrastrame, 470	Latidine, 97, 108
	Lyaconine, 224
Hydroquinicino, 425	Lyacontine, 202, 222
Hydroquimine, 424, 410, 415	Lycoctonine, 224
Hyoscine, 244, 250	
reactions of, 254	MATE, 526
saponification of, 244	Meconates, 839
Hyoscyamine, 244, 249	Meconarceino, 327
roactions of, 254	Mesome acid, 886
saponification of, 244	detection of, 338, 358
Hyoscyamus, see Henbane	Moconin, 298, 335
Нурпопо, 68	Moconidine, 294, 301, 308, 326
Hyguno, 289	Mcconosin, 336
T	Metaphenylonedianine, 86
IMIDAZOIC ACID, 24	Motatolnidine, 52
Immisorble solvents, boliavious of alka	Methacetin, 68, 85
louls with, 159	Mothocodeine, 167, 296, 324
bohavious of organic sub	Mothyl chlouds, manufacture of, 16
stances with, 158	Methyl-acotamhde, 68, 71
extraction by, 154	Mothyl alloxantin, 494
Inflatin, 196	Methy lamine, 9
Iodol, 114	- reactions of, 10
Judohuo, 115	
Isodmidine, 60	Methylamimo, 63, 71
Isotropine, 270	
T. D. CO. 1000 000 000	paranitioso, 74
JAPACONITINE, 202, 204, 220	Methyldiphenylamino, 79
benzoyl , 221	Mothylphenacotm, 84
saponification of, 204, 221, 233	Metaxylidines, 57, 59
Javanine, 892	Monammes, 3
KAIRINE, 120	characters of, 8
Karreline, 119	- distinction of, 7
Kakotelin, 388	separation of, 5
Kols, 554	Morphine, 294, 309, 326
	assay of opium for, 342
Kynurine, 168	colour-renotions of, 302, 805, 813
LANTHOPINE, 294, 301, 304, 308, 325	constitution of, 167, 298, 311
Landanine, 294, 301, 304, 308, 325	detection of, 318
Landanoune, 294, 301, 304, 309, 325	poisoning by, 350
Landanum, 850	- proportion of, in opium, 838
Lobelia, alkaloids of, 195	salts of, 311
	I man oil orr

579

Morphine, separation of, 305	Opnum, tracture of, 350
solubilities of, 301, 310	Opuum alkaloida, 293, 333
toxicology of, 855	colour-reactions of, 302
Morphiometry, 342	constitution of, 167, 296
Murexide, 480	proportions of, in opium, 333
Murexoin, 480	separation of, 305
Myoctonine, 202, 222, 225	tabular list of, 294
	poisonous characters of, 294
NAPRTHYLAMINES, 90	Otevin, 122
Naphthylamine-sulphonic acids, 9	Orthine, 29
Naphthylene diamines, 93	Orthotolnidino, 52
Narconne, 294, 301, 302, 303, 326	Ortho yldines, 57, 59
constitution of, 299	Osazones, 30
determination of, 305, 306	Oxidation colour reactions, 149, 302
Narcotine, 294, 298, 301, 302, 327	314, 868, 469, 480
constitution of, 167, 298	
determination of, 305	Oxyacanthine, 465
Neurine, 19	Ovydimorphine, see Pseudomorphine Ovyhydrastinine, 470
Nicotine, 179	
	Oxynarcotine, 294, 329
constitution of, 164	D
determination of, 161, 170, 182	Papavering, 294, 301, 302, 304, 306
— posening by, 188	329
Nitianilines, 50, 63	constitution of, 168, 299
Nitrobenzene, recognition of, 67 Nitrosamines, formation of, 7, 74	Papeverosine, 294, 329
	Parabiomacetanilide, 68, 71
Nitroso-antipyrme, S4	Paraguay tea, 526
-dimethylaniline, 75	Paranilme, 63
Nitrons and, action on monamines, 7	Paraxylidine, 57
Nonnarcotine, 298	Paregovič clixir, 353
Nuv vomica, 384	Parvoline, 97
assay of, 385	Paytamine, 392
preparations of, 386	Paytine, 392
	Piazine, constitution of, 96
ORTHOTOLUIDINE, 90	Picoline, 97, 107
Opiaume, 829	Pieraconitine, 202, 204, 221
Opiates, composition of various, 357	Pieramide, 51
Optonin, 886	Pilocaipane, synthesis of, 166
Opium, 832	Piperazidine or Pipcia/me, 106
action of solvents on, 339	Piperidine, 106, 164
adultorations of, 340	Piperine, 183, 164
alkaloids of, 293, 333	Pituine, 194
assay of, for morphine, 342	Phenacetin, 68, 81
composition of, 333	Phenacoll, 68
detection of, 858	Phenethidines, 81
- extract of, 350	Phenanthridme, 126
poisoning by, 355	
	Phenazone, 32

580 INDEX

Porsoning by toluylene diamines, 88 Phonylamine, see Amiline vermu-killers, 380 Phenyl-anthne, 73, 79 Phenyl-carbamine, 46 Porphyline, 487 Porphyroxine, 296, 330, 335 Phonyl-dimethylpyrazolone, 32 Phenylene-diamines, 63, 86 Precruitants, general, for alkaloids, 134, 158 Phenyl-hydiazides, 28 Phenyl hydrazine, 27 Propylamine, 12 Phenyl-methylpyrazolone, 31 Protopue, 296, 301, 304, 330 Pseudaconine, 202, 204, 219 Phenyl-pyrazolone, 31 Pseudacomtine, 202, 204, 216 Phenyl-methane, 68, 72 Poisoning by acetamilide, 71 --- auhydro-, 205 ---- sanonification of, 204, 218, 231 --- acoustine, 209, 236 Pseudocodeine, 323 - andine, 44, 46 Psendomorphine, 294, 301, 302, 330 --- antipyimo, 37 - constitution of, 298 - anomorphine, 320 -- atropine, 248, 261 Pseudotropue, 244, 247 - benzoyl, 244, 287 - berbeime, 462 - bi nome, 382 Paccane, 296 --- cocame, 274 Pyrazme, 30, 95 - cocamine, 286 Pv1azolc. 30, 96 --- eoderne, 322 Pyrazolines, 30 --- contonues, 174 Pyrazolones, 30 --- conine, 175 Pyndine, 96, 99 --- eurare, 388 - assay of commercial, 104 - ethylstryohnum, 19 — bases, 96 --- hemlock, 175 -carboxyhe acids, 110, 165 --- hydracetan, 28 - detection of, 101 --- bydrazine, 23 - homologues of, 107 --- hyoseyamus, 261, 250 --- walts of, 101. - laudaniue, 325 Pyridone, 96 --- lobeha, 195 Pyrodue, 28 --- Iupinme, 178 Pyrone, 96 --- metaphenylenediamine, 87 Pviiol, 96, 113 - mothagetin, 85 - methyl , 114 tetrasodo , 114 --- meoime, 183 --- opiates, 357 Pyrroline, 96 --- opium, 355 --- opium bases, 294 QUINALDING, 115 --- paraphenylenediamine, 87 Qumamicine, 427 --- phenacetan, 83 Quinamidine, 392, 427 --- protopine, \$80 Quinamine, 892, 397 --- pyridine, 98, 108 ---- constitution of, 169 - sparteine, 197 Quinazoline, 115 --- spigelme, 198 Quinetum, 448 - strychnine, 372 Quinicine, 433 - thehame, 331 Quinidine, 393, 425 - tobacco, 183 - determination of, 426, 459

INDEX. 581

Quantime, reactions of, 397, 426	Stranomum, 268	
Qumine, 393	Staychnine, 361	
constitution of, 168	- assay of nux vomica for, 385	
decomposition-products of, 168	- constitution of, 168	
determination of, in back, 449, 455	- detectant of 374	
- distinction of, from allied bases,		
405		
- formation from cupreme, 169, 398	oxidation test for, 368, 469	
hydro-, 424	poisoning by, 872	
- iodosulphate of, 403, 454	- preparations of, 376	
non cataste, 421	ptomaine simulating, 371, 375	
precipitation of, as herepathite,	reactions of, 364	
408, 454	separation of, from brucine, 366	
teactions of, 400	toxicology of, 372	
salts of, 406	vermm-killers containing, 378	
- sulphate, 406	Strychnos nua vonuca, 384	
examination of, 408	Stylophonne, 296	
impunties in, 408	Sulphamlic acid, 49	
- synthetical isomers of, 169		
- tannate of, 420	TAL B 1818, 39	
trhotme of, 423	Tua, 499	
wine of, 424	- adulterations of, 509	
Quinoidine, 488	alkaloid m, 419, 492, 504	
nodosulphate of, 454	Arabian, 527	
Qumoline, 114, 116	ash of, 511	
- tetrahydro, 119	Assam, 506	
	Bush, 503	
RESOFTRIN, 37	Cape, 509	
Rhosdine, 294, 801, 831	caper, 520	
Rubidine, 97	catechu m, 519	
	Ceylon, 506, 512	
SANGUINARINE, 295	China, 512	
Saliem, colour reactions of, 146, 147,	chlorophyll m, 505	
870, 409	composition of, 501	
Salipyrin, 37	essential oil of, 501	
Santonin, colom-resctions of, 148, 370	exhausted leaves m, 513	
Scopolamine, 244, 251	extract of, 505	
Scopoletan, 262	facing of, 522	
Smapine, 133, 167	- toreign leaves in, 522	
Snuff, 193	Indian, 503, 512	
Solanine, occurrence of, 262	infusion of, 505	
leactions of, 146	Japanese, 502, 506	
Solvents, action of, on opium, 339	Java, 512	
on plant-constituents, 151	leaves, recognition of, 523	
immiscible, use of, 154	he-, 520	
Spartemo, 197	mineral adulterants of, 510	
Spigeline, 198	moisture in, 504	
1.0		

Ten, Natal, 508, 512	Tobacco, personing by, 183
Paraguay, 526	smoke, composition of, 192
preparation of, 499	Tolidine, 90
sloe leaves in, 502, 505	Toluidino, commercial, 54
tannın ın, 515	density of, 56
tasting, 507	oxalates, 55
Trobizoudo, 527	phosphates, 51
Tannin, in collec, 529	Tolundines, 41, 51
- m tes, 491, 515	Toluylene-diamines, 87
reactions of alkaloids with, 135	Tramidophenol, 85
Tetra alkylated ammoniums, 18	Tranethylamine, 12
Totrahydro beta naphthylanum, 92	hydrochloude, 16
Totraiodo pyriol, 114	reactions of, 10
Totathyl ammonum compounds, 19,	Tuphenylamino, 80
Thalleroquin reaction, 496, 397, 401	Tuphenyhosamhuc, 61, 66
Thalline, 120	Tritopane, 294, 801, 382
cthyl , 121	Tropemes, 243
Thobame, 294, 331	artificial, 253
colour-reactions of, 302, 331	sepondication of, 241
- constitution of, 167, 296	Tropine, constitution of, 155, 246
determination of, 306, 307, 382	benzoyl-, 253
solubilities of, 301	properties of, 216
Theoliomino, 492	pseudo , 247
characters of, 493	salıcyl-, 253
constitution of, 492	Truxiline, 271, 281
determination of, 494	
in tes, 489	Unio Acib, colour-reaction of, 480
proportion of, in cocos, 496, 558,	
560	VLEAURING, constitution of, 183, 166
Thoophylline, 498	Vermu-killers, 378
Thormfugm, 122	Vinasses, 13
Thermue, 92	Vnidine, 97
Tineture of camphor, compound, 358	P
acourto, 229	Wine of quanine, 424
belladonna, 266	
conium, 177	XANTHINE, constitution of, 475
heinlock, 177	colour reaction of, 481
henbane, 268	dimethyl-, 473, 492
nux voinica, 357	derivatives of, 478
opuun, 350	- isolation of, from tea, 473
quinne, 423	tamethyl-, 478, 474
Tobacco, 184	Xanthopuccme, 471
ash of, 188, 188, 189, 190	Xenylamino, 63
combustibility of, 190	Xylıdınes, 57, 63
composition of, 184	_
extract, 193	YERBA, 526
nitrogen 10, 187, 189, 190	Yopan, 527

ADDENDA.*

Page 4. Separation of Methylammes M Delépine, Compt. rend., CXII 1064, 1272, abst. Jour. Chem. Sec., 1xx 1. 519, 588, 1xxu. 1. 585, Chim. Phys., 1806, p. 439

Page 9. Coloni reactions of Aromatic Ammes with lead oxide C Lauth, Comot rend., ext. 975, abst JCS, lx 433

Page 9 Analysis of mrytures of Ammonia and Mothylamines H Quantin, Compt ions, exv 561, abst JCS, lxiv 104

Page 9. Proparation of Ethylamine and Methylamine Tillat and Fayollat, Bull Soc Cham, at 23, abst Pharm Joun, vary 621
Page 9 Preparation of Methylamine Biochet and Cambier, J. Pharm et Cham. 1896, in 172, abst J 08, bx 1

Fage 12. Proparation of Trimethylamine F Chancel, Compt. 1880, exiv. 758, abst. J.C.S., Ixii ii 804, Bull. Soc. Chim., [3], vii 405, abst. J.C.S., Ixiv. ii 249

Page 15 Separation of Trimethylamino from Ammonia H Fleck, Amer Chem Jour., xvin. 670, abst J CS lxxii n 168 Page 22. Manufacture of Hydiazam Jour So. Chem Ind., xi 370

Page 22 Determination of Hydrazine II of mannaud Kuspert, Berichte, xxxi 64, abst Analyst, 1898, xxii 95

Page 28. Products of oxidation of Phenyl-hydraane with Folling's solution Strach and Kitt, Monatsh, xii 316, abst. JCS, lxii it. 1322

Page 28 Determination of Phenyl-hydrazme H Causse, Bull. Soc. Chim, alx 147; abst Analyst, 1898, xxiii 95

Page 28 A colour-reaction of Phonyl-hydrazine L. Simon, Compt. rend., cxxvi. 483, abst. Analysi, 1898, xxiii 131

Page 30 Use of Phenyl hybianne for the detarmination and differentiation of Sugars. Maquenne, Compt sende, cut 1990, acts. L 76 S, Ix. 1142.
Page 32 Determination of Authyrime. O Kippenberger, 2cts. acad.
Chem., xxvv 693, abst. Analyst. 1897, xxv. 219. M. F. Sobsak, Amer. Jour. 693, abst. Analyst. 1897, xxv. 219.
M. F. Sobsak,
Amer. Jour. Flan. p., Ixvv. 221, 631, abst. J.S. 61, 1896, xxv.

Page 58 Determination of Pau-tolundine. G A. Schoen, Zeit anal Chem., XXIX. 86, abst. J C S, lvin 889.

Page 54 Separation and determination of Anthro and Tolumbue Dobriner and Schianz, Zeit. anal. Chem., xxxiv. 734, abst J S C.I., 1896, xv. 298

^{*} Compiled, at the request of the Author, by Mr Arnold R Tankaid.

584 ADDENDA

Page 54. Examination of commercial Tolumbiae F F Ra a be, Chem Zeit., av. 116, abst. J S C I, v. 488

Page 57. Proporties of Panaxylidine R Michael, Berichte, xxvi. 39, abst. J C S, kxv. 1 198

Page 59. Determination of commercial Xylidines W. Vaubol, Zeit anal, Chem., xxvi. 285., abst. J.S.U.I., 1897, xvi. 639.

Page 63. Assay of Alkyl-amilines. W. Vanbel, Chem. Zeit, Vii. 485, abst. J.C.S., kny 805.

Page 64 Analysis of Audine Orls. H Reinhaidt, Chem Zeit, xvii.

413; abst. Analysi, 1893, vvin. 160.
Page 68. Micro-chemical tests for Acetsuide, Phrancelin, etc. Schoopp.

Pharm Zeat, xin 106, abst. J.S.G.I., 1897, vo. 361.
Page 68 Dotection of Acetamide in synthetical remedies, F. X. M. o. o. k.

Page 65 Properties of Phenocoll hydrochloride, Schelling, Apoll Zeat,

vi. 249, abst J S.C.I., a. 790,
Page 69 Examination of Acetambide C Plati, Jour Anal. and App.

Chem, vii. 77, abst Analyst, 1896, voi 148.

Page 69. Detection of Acetamide. F. X. M o e i k, Amer. Jour. Phan m, 1896, p. 389; abst. Analysi, va. 291. See also Analysi, 1896, λ \(\) 69.

Page 75 Action of aceta anhydride on Duncthylamine Reverdin and de la Haipe, Bull Soc Ohem, viz. 211, abst. J.C.S., lxiv i 23, Pago 81 Dulcine. See Vol III Pt. nr., page 279 tf sup

Page 82. Proparation and characters of Phenacetra, Pharm. Jour., 1889,

Page 82. A reaction for Phenacolin. Autenioith and Hinsberg, Arch. de Pharm, caxix, 456, abst. Analysi, 1892, vol. 56.

Page 83 Colour-reactions for Phonocettin, Methanetin, etc., in ninkings T G Selmi, Chem Zeil, vv. 368, abst J.S.C.I., 1893, τι. 466.

Page 83. Guastistates that Schwait's test for the detection of Acet ambde is unchable See J C S., Ivv. 432.

Page 88. Physiological action of Tolmylone dimmine. W. II unter, Jour, Pathol and Bacteriol, 1895, p. 259, abrit. J.C S., havin, in 156.

Page 89 Determination of Benedino and Tolidine W Vanbel, Zott and Chem., New 168, abot. J.C.S., Ix. u. 507.

Page 91 a-Naphthylamme has not much smell, the diagnating odom commonly sembed to it being mainly due to impunities. A page a little having a melting point of 47° C. is now made on a communical scale

Page 92. Assay of Naphthylamme sulphome acids. W. Vaubel, Chem. Zut., xvu. 1265, abst J.C.S., lxvi ii. 74

Page 92 Preparation of S Naphthylamine-sulphonic acid. A G Green, J S.C.I., 1889, vin 878, 1890, ix 934.

Page 103. Action of cadmium salts on Pyridine and Pipendine R. Varet,

Compt void, exv 464, abst J.C.S., laiv. 1 48.
Page 103. Action of tanum on Pyridine and Piperidine O. de Coninck,

Page 103. Action of familia on Pyridine and Piperidine O. de Coninck, Compt. rend., exxiv. 506, abst. J.S.C.L., 1897, xii 470.

Page 106 Piperkine and Piperazine. See also Vol. III Pt. itt., pp. 38, 194, Page 107 Properties of Piperakine Utate: Tunnie liffe and Rosenheim, Brit Med. Jour., 5th Pubnary, 1898. ADDRNDA 5

Page 107 Potassio iodide of bismuth precipitates Piperidine from its solutions or concentration and cooling, in the form of thin, transported, yellow, hevagonal plates, of characteristic mucoscopic appearance.

- Page 114 Source and properties of "Iodol" A. Trillat, Monte Secont, 1802, p. 838, abst JSCI. xi 1030
- Page 129 More chemical recognition of Alkaloids L. Elleva, Pharm Jour., vvin 48 H Barth, Pharm Jour., 1898, it. 635, 1899, 1360 Vadam, J. Pharm Chem., 1896, [4], 485, abst. J.S.C.L., 1897, xvi. 185.
- Page 180 Titation of Alkaloids L. F Keblei, Ame: Jour Phaim, layin 499, abst. Chem. News, laxin, 298 H Biown, Phaim Jour, axv, 1180.
- Page 130 Action of acids on Alkaloids A. H. Allen, Chem. News, lxvi. 259
 Page 131 Volumetrie determination of Alkaloids. E. Léger, Compt. 1 cnd.,
- exv 732, abst. J S.C I, 1893, vu 470
 Page 131 Titistion of Organic Bases with methyl orange. G Lunge,
- Chom Ind., vvi 490, abst J.S.O.I., 1894, viii 667.
 Page 186 Tannates of Alkaloubs are soluble in glycerin Thus allows of a ready separation of the alkalouds from albumin. (C Kipponberger, Analust. 1895, xx. 201)
- Page 187. G. Beitiand (abst *Phurm Jous*, 1399, t. 503) strongly recommends Silico tunes to each as a precipitant of Alkaloids
- Page 188 Delicacy of Marmé's reagent for Alkalouds. S Volven, Ann. de Pharm, ann. 145, abst. Analyst, 1897, vm 241.
- Page 138 Alkaloidal Periodides Determination of Alkaloids Prescott and Gordin, Amn. Jour Phanm, lxx 439, lxx, 14, 18, 4bst, Analyst, 1898, xxiii 324, 1899, xxiii 74, 75
- Page 189 Volumetric determination of Alkaloids L. Barthe, Compt.
- Page 139. Volumetaic determination of Alkaloids P C Plugge, Compt.
- Page 151 Determination of Alkaloids Grandval and Lajoux, J.

 Pharm et Chim, xxviii. 99, abst J.C.S., lxiv. ii 608 H A. D.
 Jowett, Pharm Jour., 1899, i. 377.
- Fage 151 Examination of decomposed human ioniams for Alkaloids. C Kippen beiger, Zeit anal. Chem., xxxiv 294, abst J C S., 1895, level. u 485
- Page 151 Determination of Alkaloids K Dioterich, Pharm Zett, xliv. 242, abst Pharm. Jour, xxv. 962
- Page 151 Isolation and determination of Alkalouds C Kippen beinger, Zeit, anal Chem., XXV 10, 407, abst JOS, IXV ii 681, 682, Analysi, XXI 191 Fair and Wright, Pharm Jour, 1897, 1 202.
- Analysis, xxi 191 Fall and Wilgit, Fall and Jansen, 1895, 18
- Page 160. For the separation of Alkaloids in toiensic cases, C. K. 19 penber g or agitates the alkaloidal solution, first with sulphunce and and chloroform, then with caustic soids and chloroform; next with sodium brarbonate and alcoholic chloroform, and finally estimates with sodium chloride and agitates with other chloroform, which last trustment to-

moves Strophantinn. (Zeit anal Chem., 1895, p. 191, abst. Analyst, 1895, xx. 201.)

Pago 160. Claus' method of Tea assay is valueless. Compare page 186.

Pago 161. Lioyd's process is stated to give more reliable results than any other rapid mothed of Alkaloidal Assay (Nichols and Norton, Jour and and Appl Chem, vi 162, abst. J S C.I., 1863, vn. 68).

Page 173. New test for Conne Van Senus, abst. Analyst, v 79.

Page 174 Preparation and Proporties of Contestine, Commi, etc. Lell mann

and Mullor, Benichte, XXIII 680, abst. J.C.S., lvin. 802
Page 178. Volumetric determination of Communical Nicotine in the same

solution. G. Heut, Arch de Pharm, cenxxi. 376, abst J C S., INV. 608
Page 180 One c.c of normal hydrochloric or sulphure and is nontralised

by 0 162 grammo of Nicotino, when methyl-mange is used as the indicator.

Page 182. Determination of Amnionia and Nicotine in Tobacco. V

Page 182. Determination of Ammonia and Nicotine in Tobacco. V Vodrodi, Zeit and Chem., 1895, dost Analys, 1895, N. 255 R. Kissling, Zeit and Chem., NXIV 731, abst. J.S.C.I., 1896, vv 300. A Pazzoluto, abst. J.C.S., lx 771

Page 184 Analysis of the Tobacco-plint li J Davidson, abst. J C.S., law, n. 38

Page 190 Composition of Tobaccos H B Cox, Pharm Jour, xxv 550, Page 192 Composition of Tobacco smoke A. Gantiol, Compt. 100d, cvv. 992; abst. J.C.S., lvv 1 226.

Page 193 Examination of Tobacco-extinct J. Pineito, Chim. Zell., xvi 178, Analyst, 1892, xvii 178.

Page 195 Timeture of Lobelia J F. Liveiseege, Pharm. Jour., lv 141.

Page 108 For papers on the Archite Alkdovie by Dunstan, Tunney, Dunstan and Cair, etc., wo Jear. Chem. Soc., iv., 385, 361; iv.it., 448, 401, 401, 1vr. 178, 290, 1vr. 1908, 1vr. x. 1921, 1vr. x. 293, 1vr. 306, 30rr. Sc. Chem. Led. xi 306, Physm. Joney, xxii., 458, xxii., 85, 625, 765, 1046, vr. vr. 561, 739, 736, 561, 910, 645, xxv. 778, 860, 928, 1117, 1896, 1, 121; 1896, 1, 205.

Page 226. The extraction, composition, and properties of Atisine and its salts H A D Jowett, J.C.Y. 1818.

Page 244 and 250. O Hease has shown that Hyoseme probably has the formula C_BH_BNO_B, and by superincation yields the base oscina C_BH_BNO_B and not proud otroping (Ann. dos Chemes, colan. 100; abst. Phorm. Join. [3], von 221)

Page 247 Some new Gold Salts of the Mydnatic Alkaloids II. A D Jowett, J C S, INA 679

Pago 251. For information to pecting Apo-akopane, Attopanuac, Bella donne and Scopolanuac, see papers by R. Schmidt and O. Hesse, abst. Pharm. Jour., [3]. xxii. 1021; xxii. 221, [4], 1890, 1, 383.

Page 254. Soparation of Attopino and Hyosoyamine O Hosse, abst Phain, Jour., [3], xxiii, 201

Page 258 Test for distinguishing Atropine from Strychime D Vitali,

ADDENDA 587

Chem Contr., 1804, n. 816; abst. J C S., 1895, lxvni n. 467, Zed anal. Chem., xxxvni. 184; abst. J S C J., 1890, xvni. 404.

Page 261 Toxicological detection of Atopino and its allies Ciotto and Spice, abst. J.C.S., 1x 772.

Page 261 Detection of Attorne in forensic cases P Solstein, abst Analyst, 1897, xxii 162.

Page 264. Assay of Belladonna W A. Puckner, abst. Pharm. Jour, 1898, 11 97

Page 266. Assay of Belladonna plasters. C E. Smith, Amer Jour. Phasm., lxx. 182.

Page 266 Assay of hqmd exhact of Belladonna H Wilson, Pharm Jour, 1898, 1 450 E Dowlaid, Chemist and Druggist, 1899, p. 401 F. C J. Bild, Pharm Jour, 1899, 1 432

Page 272. Isolation of Cocame from accompanying alkaloids. Einhoin and Willstatter, abst JCS, lxvi. 1. 478

Page 274 Test for Cocame Scharges, Chem Cents., 1893, n. 888, abst. J C.S., lxvi n. 127.

Puge 274 Properties of Engane and Cocanne, G. Vulprus, alast JSCI, 1896, xv 679, 745 P Silex, abst JSCI, 1897, xv: 631. Page 274 Reactions of Cocaine JC Stead, Pharm Jose , xxi: 002. A

Kubsine, Pharm Cents, axxiii 411, abst JSCI, 1894, au 380
Page 274 Detection of Cocame in poisoning cases. H W Glascia, Chem Cents., 1894, ii 220, abst JUS., 1895, livin ii 336.

Page 274. For the detection of Oxame, A Kuborn o Jun (Oken News, Ivrn. 264) tecommends that I or of inthe and I/129 mg, I) bended to the substance as a pocelar dish, and the input ovaposated at 100°C. When cold, a drop of alcohole postals added No colour is produced in the cold (datameteon from anopure), but when heated on the water-bath, an intense voice coloutour is suddenly produced.

Pages 27 and 288. Reactions of Cocame and Ecgonnic. D Vitali, abst JOS, lx 1561.

Page 280. Characters of Cocame hydrochloride Paul and Cownley, Pharm Jour, 1808, 1 586

Page 280. Test for the purity of Cocame salts. G L Schafer, A J Cownley, Pharm. Jour., 1899, 1 386

Page 280. A new Alkaloid in Coca leaves. G L Schafel, abst Pharm Jour., 1899, i 350

Pago 280 Maclagan's ammonia test for the purity of Cocaine hydrochiloride. See absts. Pharm. Jour., 1898, 1 449, 473, 1898, u. 26, 1809, 1 431

Page 287 Properties of Benzoyl-pseudotropine and its selts Pharm Jours., [3], axin. 241

Page 293 Assay of fluid extract of Coca. C T Kingsley, Amer. Jour Pharm, 1896, p. 609, abst Analyst, 1897, xxii 77

Page 293 The author was indebted to Mi D B Dott for pousal and contection of the section on Opinin Alkaloids.

Page 295 Assay of Sangunaria and its preparations C H. La Wall, Amer Jone Pharm, 1896, p. 305

Page 295. Reactions of Chelidonius with phonols J. A. Battandier, Compt. 1end, exx 270, abst. J.C.S., 1895, lyvin. ii. 886.

Page 300 Solubility of Morphine and Narcotine. E. L. Patch, Amer Jour Pharm, 1808, p 553

Page 305. Detection of Alkaloids by the Stas-Otto method. R Otto,

Arch de Pharm, cryxxiv 317, abst JCS, lvx. 11 508
Page 312 Denvatives of Morphine (Morch's Report, 1898) Pharm Zoul

xlv., 117; abst JSCJ, 1899, xvii 395
Page 312 Proportios of Diomine L. Hosse, Pharm. Centr., vi. 1, abst.
Analysis, 1899, xxiv 198.

Page 318. Colour tests for Morphine G. Bruylants, J. Pharm. et Chim, May 1st, 1895, abst Pharm John., xxv, 1123.

Page 313 Reactions for Morphine, G Bruylants, Bull. Soc. Chim, xm 497, abst. J.C K., lx n 132.

Page 315 The Furfural reactions of Alkalouds. N. Wender, Chem Zent,

vvi 550, shat J.S.C.I., 1393, an 369
Page 316. The determation of Morphuse. O Kippenberger, Zect.
and Chem. vvv 421, abst Analyst, axii, 42,

Page 816 The determination of Alkalous in Naicotar extracts. J. II Schmidt, Chem. Zeit, xvi. 1275, abst. J. S. C. I., 1893, xii. 470

Page 316 Titration of Morphine Unineprin and van Erjk, Eadl. So. Chim, 1x 437, abst J C.S., lvv. n. 607.
Page 317 Fourtewards test for Morphine School, Arch. de Pharm.

Page 317 Funityando test for Morphine Schaet, Men. de Pharm oxxxiv 348, abst. Pharm Jour, 1890, n. 61.

Page 321 Branmaton of Codeme Tambach and Honke, Pharm

Unite, xxxviii. 150, abst Analyst, xxii 210
Page 223. Semantion of Colone and Mornings. L. Fountet. J. Pharm.

et Chem., xvii 49, abst JSCI, 1897, xvi 159 Page 327 Reactions for Navenne and Papaverine. C Kippenberger,

Zett anal Chem , 1896, p. 294 , abst Analyst, 1895, \(\cdot 201. \)
Page 331. The chemistry of Thebane. M Fround, Beruhle, 1897, p. 11 ,

alst J C S, Lvin L 117; lvn 1 494
Page 840 Determination of starch and strontium sulphate in Opium.
Kebler and La W all, Ame. Jon. Pharm. 1897, p. 214.

Page 342. Assay of Opum D B. Doti, Phum Jonn., [3], vev. 847.

Page 3-9. Assay of Opinm and the propagations. O in und wit land Layoua, J. Phenn. et Chim., 1867, p. 150, abst. J. Sc. C., 1867, xv. 265 G. Looff, Apach. Zerl., 1869, n. 192; abst. Analysis, vo. 186. F. X. Mootk, Anne Jose, Pages ann., 1867, page 541. E. J. Mitland, Pharm. Jone, pages 831. D. B. Dott, Pharm. Jone, 1892, p. 746; 1894, p. 747.

Page 351 Assay of Laudanum. L F Kobler, Amer Jour Pharm., 1893, p 209

Page 356 Tests for Morphine in forence cases D. L. Davoll, Jun., Ame. Chem. Jour., xvi 790, abst. Analyst, xx 38 J. B. Nagel voort, Amer. Jour. Pharm., 1806, p. 374

Page 363. Liquor Strychnuse Hydrochlondt (BP, 1898). Martindale,
 Lunen, and others, Pharm Join, 1898, 1, 587; 1898, 11, 19, 43,
 67, 1899, 1, 190.
 Page 369.
 Weter of contribution of Strucking Habridge 19, 12

Page 363 Water of crystallisation of Suychnino Hydrochloride. D. B. Dott, Phurm. Jon., 1899, 1 53 W. H. Martindale, and p. 120. ADDRNDA 589

Page 364. Alkaloidal content of Strychnine salts W. Duncan, G. Coull, Pharm Jour, xxii 843, 846

- Page 864 According to D. B Dott (Pharm. Jour, [3], xxm. 197), the
- solubility of Strychnino Hydrochlorule in cold water is 1 in 35
 Page 864. Detection of Strychnine in forensic cases. A. S. Ou shiman,
- Chem. Centr., 1894, n. 461, abst. J C.S., 1895, lavni n. 542
 Page 367. The following Alkaloids are not precipitated by potassium feito-
- cyanide —attopine, codeine, emetine, narceme, spatteine, "veratine"
 Page 368. Action of sulphune and on Strychnine. Bailey and Lange,
 Amer Jour Pharm., 1898, p 18
- Page 368 Examination of the Oxulation test for Stryohnine Mason and
- Bowman, Amer Chem. Jour, xvi 824, abst JS C.I., 1895, xv 313.
 Page 308. Detection of Stayehnne H Bockurts, Arch de Pharm, abst Pharm Jour, xviv. 2
- Page 383 Colour-teactions of Bucine. P Pichard, Compt and , exam. 590', abst Analyst, 1897, van 47
- Page 384 The Brunne and Strychnine in mix vomica seeds exist in separate cells Sauvan, J. Pharm. et Chim., vi. 1.497, abst. Pharm. Jou., xxv 1090
- Page 386 Determination of Nux Vonnea Aikalouls, C C Keller, Apoth Zett., vm 542, abst. JSCL, 1894, xm 1105.
- Page 388 Most specimens of Curare contain methyl strychime, which is one of the most active ingredients See E Auquetil, abst Pharm. Jun., xxiii. 624.
- Page 396. Test for Cinchona Alkaloids Jawo 1 owski, J. Pharm et Chim. 1896, page 553, abst J.C.S., lv. 11 629.
- Page 401 Modifications of the Thalletoquin reaction J. Due ommon, Chem Zeit, 1895, p. 214, abst. Analyst, xv. 234 F. S. Hyde, Amer. Chem. Low. 1891, 1891, abst. Analyst, 1897, xvii 266
- Chem Jour., xix. 331, abst. Analysi, 1897, xxii 266
 Page 401. A reaction for Quinne. C. Cailez, J. Pharm et Chim, 1896,
- p 253, abst JOS, lax n 584
 Page 402. Determination of Quinnie L Baithe, Compt rend, exv
 1085, abst. JS, UI, 1893, xii 380
- Page 403 Titration of Qmmme L F Kebler, Amer Chem Jour, 1895, xvn 892, abst J.O.S., lxx n 551 A. H Allen, Analyst, 1896, xxi 85
- Page 408. The basetty of Qumme Howard and Howard, Pharms Joss, 1898, 1 154.
- Page 408. The testing of Quinine Sulphate. M. Knbli, Chem Centr, 1895, n. 1058, abst. J C.S., lxv n 550, lvvn n 168 O Hosso, Arch de Pharm, cexxiv. 195, abst. J.C.S., lxx n 550
- Page 408. A test for the purity of Qumine Salts J. de Vrij; abst. J.S C I, 1897, xvi 165
- Page 408. Tests for Quinne. T. G. Woimley, Amer. Jour Pharm., lxvi 561, abst. Pharm Jour., xxv 542.
- Page 446. Assay of lineture of Cinchona. Fair and Wright, Pharm Jour., xxin. 248
- Page 445 Valuation of Cinchona Extract, M. L. Hulsebosch, Chem. Centr., 1896, 1, 141, abst. J.C.S., lxx n 682

590 ADDENDA

Page 449 Determination of the Alkalouki u Cuchona Balk. M. I. III lishobach, Phaum. Gair. v. v. 229, Jubr 4. Z. G. I. 1869, x. 537. W. Hatubansach, Phaum. Gair., v. xxu. 294, abrt J. S. C. I. 1869, x. 57. 79 J. H. Schmidt, Chem Zeit, v. v. 307, abrt J. S. C. I., 1892, v. 779 J. H. Schmidt, Chem Zeit, v. v. 307, abrt J. S. C. I., 1899, xvu. 408. H. Nika no. 4. dir Mer Pharm. 1898, p. 328; abst. Amm. Jon. Thurm. April. 1899

Page 467 Constitution of Hydiastine and its derivatives Fritsch, Liebig's Annalen, colxxxvi, 21, abst Pharm Jons, xxv 1193

Page 467 Properties of the Alkaloids of Hydrastis Genadenses K. von Bunge, Chem Centr., 1895, 1 1173; abst. J.C.S., lxx is 492.

Bunge, Chem Crar, 1983, 1 1173; and J. G.S., 188 ii 402.
Page 467 The Chemistry of Hydrastac and its salts. Freund and Dormeyer, Berichle, xxiv. 2730, 3164, abst. J. C.S., ix. 1518, iv., 1 223

Page 468 Reactions of Hydrastine and other alkaloids D. Vitali, abst J C S. lvn 1, 756.

Page 489 Determination of Caffeine E Tassily, Bull. Soc. Chim., xvii 598, 708, 761, abrd J.S.C. I, 1897, vii 697, 831 M Gombierg, Amer. Chem. Jour., viii 331, abrd. Analyst, 1896, vvi. 193 A Delaconi, J. Pharm. et Chim., iv 490, abrd. Analyst, 1897, vvii 76.

Page 489 Debermantono (Caffence For acter and Reverbell us un, and Hilger and Juck Onno E. (Mens. Cett., 1987, [1], 757; abed. Analyst, 1597, xu. 158, 238 G. L. Sponeer, Anno Ichen. Jour., xu. 192, abet J. C. N., 1514, 901 G. C. K. Filler, (Mens. Zeut., xu. 102, abet J. S. C. L. 1897, vor. 1988 W. A. Puck no., Amer. Chem. Jour., vor. 198, abet. J. S. C. J. 1898, xv. 292. Pettic and Tart., Bull Soc. Chem., vol. 85, D. J. 1898, xv. 292. Pettic and Tart., Bull Soc. Chem., vol. 185, abet. Analyst, 1898, vol. 292. F. V. H. C. Chem., vol. 185, abet. Analyst, 1898, vol. 292. F. V. H. C. Chem., vol. 185, abet. Analyst, 1898, vol. 292. Pettic and Tart. 1998, vol. 293. Pettic and Tart. 1998, vol. 293. Pettic and Tart. 1998, vol. 294. V. H. C. Chem., vol. 294. Analyst, 1898, vol. 1915, abet. J. J. S. V. V. L. C. Chem., vol. 295. C. C. L. V. V. L. N. V. L. V. S. V. V. 1808, Chem. 1918, vol. 294. V. V. L. V. S. V. V. 1809, C. H. La Wall, Amer. Jour. Pharm., 1897, p. 350, abet. Analyst, 1897, xan 298.

P.199 490. Determination of Caffeine in Tea. N V Sokoloff, abst. JUS, Juv 1. 362. Kollner, Forechings berichte, w 88; abst. Pharm Jour. 1897. n. 83

Page 860. Determination of Caffeine in Ton. R. II Gane Glows. See Chem. Ind., 1880, page 99 states that, after a tail of swend pacessage, he bound the anthor's method to give the best and most concentent insents. A comparison of the isentile obtained by Gane with the methods of Paul and Go will be you do of the author shows that the latter process gives intended to higher penish of suffuent that the form, whilst the althody and obtained in a state of great purity. Gase regards the author's unbold as the to first printing the control of the

ADDENDA, 591

Page 493 The Examination of Theobiomine. M Figure 53, J Pharm et Chim., vn. 521, abst Analyst, 1898, xxii, 213

Page 498 Theobromme and its homologues. Brunner, and Binnner and Leins, abst. JSC, I, 1898, vvi 78, 946.

Page 496. Determination and separation of the Alkaloids of Cocca. Brunner and Leins, Chom. Cents, 1898, p 512; abst J S.O.I., 1898, xvn 991.

Fago 498. Determination of Theobromine in Cocca, etc. Hilger and Emingen, Forest. Ber., 1895, p. 299, abst. J.C.S., leviu 542. L. Maupy, J. Pharma. Chem., 1897, v. 329, abst. J.S.C.I., xv. 041 P. Suss, Zet. and Chem., v.xxi. 57, abst. J.C.S., klv. 198

Page 496. W. E. Kunze (Zest anal Ohem, xxxiii 1, abst Analyst, 1894, page 194), has proposed the following method for the determination and

separation of the Åltaladus of Coca —
For the estimation of the total alkaladus, ten grammes of the cocas is builded for twenty mustus with about 150 c o of five pet cent, sulphune cond. Altered, and the results throughly washed with boiling wats. The alkaladus are prespirated from the fittate by a large excess of a nutra cand outbroot of solumn phophomoshybots, and the luqual Kept warm for twenty-four hours. It is then illiered, the precipitate washed with the dilute sulphune and, and at once decomposed by baya-water, the excess of harum being precipitated by passeng curbon diovide thiooght the luquid. The builded simple progritists are together originated to dryness, duried, and tone, then all the distributions of the simple control of the cont

For the separation of the caffane and theoloronus time obtained, so not form a similar computed into its monthlos aires aid. (Caffane and control form a similar compound.) For this purpose, the mixed alkalonks are disvoled in ammonia, a considerable eccess of aires mixed aid, and the liquid boiled downs a every small built, and until all flow ammonia verxysiled. The crystalline pencipitate of theoloronus-civer and (CHA-gX/G_o) is collected, well weaked with boiling water, dried, agented, and the resultant view weighted 11 fla known measure of standard stirve mixeds be employed, the amonut of theoloronus presuptated may be deduced from the occess of writer contained in the filtrate as determined by Othmat's suchod After the tatation, the alkalonks may be readily extracted from the precipitate and dilutes, and tested as to them purply, extracted from the precipitate and dilutes, and tested as to them purply, extracted from the precipitate

Kunze's paper contains a valuable reamé and entreism of the methods hitherto employed for the separation of the cocoa alkalous, and the substantial accuracy of his process is confirmed by analytical data.

Page 504 Analysis of Tea Domerche and Nicolas. J Pharm. et

Page 504 Abalysis of Tea Domergne and Nicolas, J Phalm. e Chim, xxv 302; abst J C S., lvn n 926

Page 506 Detection of Extracted Tea W. A. Tichomilow, abst. Chem. News, lxvii, 196

Page 509. New Adults ant of Tea Delaite and Lonay, Bull A Belge Cham, x1 13, abst. JSC.I, 1897, xv1 700.

Page 516 The analysis of China Teas P. Dvorkovitz, abst. Jour Soc Chem Ind., x, 276.

- Page 518. In employing Edici's process for the dotermination of Taimin in Tea, the excess of copper may be determined by feirogramide. Maltacheffsky, J. Theom. Cham, xxii 270, abst. J.C.S., ix 132
 - Page 520 The composition of Caper Tea. C. Esteouit, Analysi, xxv 30. J White, 4nd, p 117.
- Page 526 The composition of Mate or Paraguay Tea H. Kunr Krause, Arch de Pharm, crxxxx 613, abst Pharm Jou., xxv 442, McKendrick and Harris, Pharm Joss., 1898, ii 52
- Page 526 Contabutions to the study of Mats. P. Macquaire, J. Pharm et Chem, 1896, p. 346, abst Analyst, xxn. 18 B. A. Katr, Zeil Nahr Untersuch., x 364, abst Analyst, 1897, xxh. 41.
- Page 526. The word "mate" is need adjectively, referring to the gourd from which the scalding infraion is sucked through the hombilla, that is, a tube having a bulb at one end. We should, therefore, always
- say "Yenba Mate," the gound plant
 Page 527. The composition of Catha edulis E Collin, Phanm Jour,
 xxiv 345
- Page 527 The following analyses of "Coffee Tea" (coffee leaves) are from the Lancet, 5th August, 1898 -

	Whole Lonf	Small Broken Lea
Cafferne,	2.66	3 20
Tannın,	7 14	6.66
Extract,	39 45	34 40
Moisture,	7.60	7.69
Ash.	8 10	5.50

- Page 528 Proportion of various constituents of Coffee. Herfoldt and Stutzel, Zeil. angew. Chem., 1895, p. 469, abet J.C.S., ix u. 63. Page 528 Proportion of water in law Coffee. ii Niederstadt, Forsch
- Ber, 1897, p. 141, alist Analyst, 1897, von. 322
 Page 528 A new Alkaloid of Coffice (Cofficience) Forster and Reachel
 mann, Pharm Zett, vin 309, abst Pharm Jour, 1897, n. 84. P
- Paladino, abst. Analyst, 1895, xx. 141
 Page 530 Alteration in composition of Coffee during reaching Hilger and Juckenack, Forch. Be, iv. 119, abst. Analyst, 1897, AM
 287. H Jacok le, Zeit f Untersuch, 1898, p 457, abst. Analyst, 1898, xxii. 244.
- Page 53 The Carbohydiates of the Coffee benry E E Ewell, Amer.

 Chem Jour, xiv 473, abst JSCI, 1393, xii 611
- Page 533 A Ptomaine in Coffee, S. Belin, Zit angew. Chem., 1898, p. 658, abst. Analyst, 1899, xxiv 36.
 Page 534. Glazed Coffee benies. E. Hanansek, abst. Analyst, viv 38
- Page 535. Exhausted Coffee benies P E Hamel Roos, abst Analyst, xxv 36 xvi 160
- Pago 535 Analysis of a spinious roasted Coffee M Maljean, J. Pharm. of Chim., 1896, p. 352, abst. Analyst; 1897, xxii 17
- Page 588. Adulterations of Coffee G. Wilti, Zeit f Untersuch., 1898, p 248, abst Analyst, xviii 209 Ponimain and Moor, Analyst, xx. 176

ADDENDA.

Page 538 Variations in the composition of Chicory. B Dyer, Analysis, 593

Page 538 Sugar in iosated Clincoly E. G. Clayton, Analysis, vs. 12 Tage 559. Determination of Caranel in Coffee tousted with sugar

Flesenius and Glanhut, Zest anal. Chem., XXVI 225; Rost.

Page 559. The analysis of Chicory. A. Ruffin, Chem Centr., 1898, p 1147.

Pago 544 Analysa of Dandelson 100t (Taravacum) L. E Sayre, Amo. Page 545 Composition of the Ash of Coffee F W Dafert, abst. J C.S.

Page 547. Analyses of "coffce-palace" Coffce Infusions E G Clayton,

Page 554 Composition of Kola-nuts. Uffelmann and Bonies, Zed

angere Chem, vxiii 710, abst Analysi, 1805, xx 42 Knov and Plescott, Amer Chem Jour, x. 34, alast Analysi, 1897, Nr. 1dl. E Knebel, Apolt Zett, vn 112, abst J S. C I, 1892, vt 545 R. Dieterich, Apods. Zeut., M. 810, abut JSOI, 1897, XVI 180 P. Carles, J. Phanm is Chim, xvi. 104, Ann der Chamie, Alli

Page 554. False Kola nuts. J. H. Hart, Pharm. Jour., 1898, 1 184 Page 554 Kolann and Coconn, the Gincorules of the Kola-unt C

Solweitzei, Phasm Zest., xhn 380, 389, abst Phasm Joss., Page 554 Determination of the Alkaloids of Fluid Extract of Kola O

Schumm, Apoth. Zest., 1898, p 682, abst J S C I., 1899, xviii 408. Page 558. Determination of the chief constituents of Cocca Boans II Bookurts, Avib de Pharm, CEXXI. 887, abst J T.S., IVI II 363 Page 559. The statch of Cocoa. E. S. Bastin, Ams. Jour. Pharm., Lvn.

Pages 561 and 566. Sugar in commercial Cocoss M. Schiodol, Zett

angew. Chem , 1892, p 173 , abst JOS, hu 100

Pages 561 and 566. Sugai in Chocolate X Rootines, Ann. der Chimic, 1898, P 288, abst Analyst, 1898, Xu 256 P Carles, J. Than m. et China, viii 245, abst JSCI., 1898, vvi 1076 H Pellet, Bull A Belge Cham, xiv. 790, abet J S.C I, 1897, xvi 474.

Page 581 Detection of Arachis in Chocolate A Bilteryst, Bull, 4 Belge Cham, x 447, alst Analyst, 1897, xxii 215 Page 562 Detection of gelatin in Chocolate P On Livy, J Pharm at

Page 583. Analyses of commercial Cocoas, J. S. Livoisedge, Pharm.

Jour., XXIII 922 F Yaple, Ames Jour Pharm, 1895, p. 318 Page 564 Determination of "soluble" constituents in Cocon. A Stutze: Zet angew. Ohem., 1892, p. 510, abst. J.S.O.I., 1893, M. 54 Page 567 Detection of milited stateli in Cocoa Posetto, Phas in Centr.,